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(54) Title: ARTEMIN, A NOVEL NEUROTROPHIC FACTOR

#### (57) Abstract

A novel growth factor, artemin, which belongs to the GDNF/neurturin/persephin family of growth factors, is disclosed. The human and mouse amino sequences have been identified. Human and mouse artemin genomic DNA sequences have been cloned and sequenced and the respective cDNA sequences identified. In addition, methods for treating degenerative conditions using artemin, methods for detecting artemin gene alterations and methods for detecting and monitoring patient levels of artemin are provided.

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WO 00/18799 PCT/US99/22604

### ARTEMIN, A NOVEL NEUROTROPHIC FACTOR

### Reference to Government Grant

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# 5 Related Applications

This application, which claims priority to Provisional Application Serial No. 60/108,148 filed November 12, 1998, is a continuation-in-part of Application Serial No. 09/163,283, filed September 29, 1998.

#### 10 Background of the Invention

### (1) Field of the Invention

This invention relates generally to trophic or growth factors and, more particularly, to a new growth factor, artemin, which is a member of the neurturin-persephin-GDNF family of growth factors.

### 15 (2) Description of the Related Art

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The development and maintenance of tissues in complex organisms requires precise control over the processes of cell proliferation, differentiation, survival and function. A major mechanism whereby these processes are controlled is through the actions of polypeptides known as "growth factors". These structurally diverse molecules act through specific cell surface receptors to produce these actions.

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Growth factors termed "neurotrophic factors" promote differentiation, maintain a mature phenotype and provide trophic support, promoting growth and survival of neurons. Neurotrophic factors reside in the nervous system or in innervated tissues. Nerve growth factor (NGF) was the first neurotrophic factor to be identified and characterized (Levi-Montalcini et al., J. Exp. Zool. 116:321, 1951). 5 NGF exists as a non-covalently bound homodimer that promotes the survival and growth of sympathetic, neural crest-derived sensory, and basal forebrain cholinergic neurons. In sympathetic neurons this substance produces neurite outgrowth in vitro and increased axonal and dendritic growth in vivo. (See Levi-Montalcini and Booker, Proc Nat'l Acad Sci 46:384-391, 1960; Johnson et al. Science 210: 916-918, 1980; 10 Crowley et al., Cell 76:1001-12, 1994). NGF has effects on cognition and neuronal plasticity, and can promote the survival of neurons that have suffered damage due to a variety of mechanical, chemical, viral, and immunological insults (Snider and Johnson, Ann Neurol 26:489-506, 1989; Hefti, J Neurobiol 25:1418-35, 1994). NGF also is known to extensively interact with the endocrine system and in immune and 15 inflammatory processes. (Reviewed in Scully and Otten, Cell Biol Int 19:459-469, 1995; Otten and Gadient, Int. J. Devl Neurosci 13:147-151, 1995). For example, NGF promotes the survival of mast cells. (Horigome et al. J Biol Chem 269:2695-2707, 1994).

In recent years it has become apparent that growth factors fall into classes, i.e. families or superfamilies based upon the similarities in their amino acid sequences. These families include, for example, the fibroblast growth factor family, the neurotrophin family and the transforming growth factor-beta (TGF- $\beta$ ) family. As an example of family member sequence similarities, TGF- $\beta$  family members have 7 canonical framework cysteine residues which identify members of this superfamily.

NGF is the prototype of such a family of growth factors. Brain-derived neurotrophic factor (BDNF), the second member of this family to be discovered, was shown to be related to NGF by virtue of the conservation of all six cysteines that form the three internal disulfides of the NGF monomer (Barde, *Prog Growth Factor Res* 2:237-248, 1990 and Liebrock et al. *Nature 341*:149-152, 1989). By utilizing the information provided by BDNF of the highly conserved portions of two factors,

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additional members (NT-3, NT-4/5) of this neurotrophin family were rapidly found by several groups (Klein, *FASEB J 8*:738-44, 1994).

Recently, a new family of neurotrophic factors has been identified whose members are not structurally related to NGF and other neurotrophins but are structurally similar to TGF-β. As described in U.S. Patent No. 5,739,307 and copending application 08/931,858, the known members of this subfamily of the TGFß superfamily include glial cell line-derived neurotrophic factor (GDNF), neurturin (NTN), and persephin (PSP). The placement of GDNF, neurturin and persephin into the same growth factor family, also referred to as the GDNF ligand family, is based on the similarities of their physical structures and biological activities. Human persephin has about 40% sequence identity and about 43% sequence conservation with human GDNF; and about 49% sequence identity and about 50% sequence conservation with human neurturin. Similarly, human neurturin has about 43% sequence identity and about 53% sequence conservation with human GDNF. In addition, these three proteins have seven cysteine residues whose positions are exactly conserved. GDNF, neurturin and persephin each support the survival of dopaminergic midbrain neurons, and spinal and facial motor neurons, in both in vitro survival and in vivo injury paradigms, identifying these ligands as potential therapeutic agents in the treatment of neurodegenerative diseases (Henderson et al., Science 266, 1062-1064, 1994; Horger et al., J Neurosci. 18, 4929-37 1998; Lin et al., Science 260, 1130-1132, 1993; Milbrandt et al., Neuron 20, 245-53, 1998; Oppenheim et al., Nature 373, 344-346, 1995), reviewed by (Grondin and Gash, *J Neurol.* 245(11 Suppl 3), 35-42, 1998). However, while GDNF and neurturin both support the survival of many peripheral neurons in culture, including sympathetic, parasympathetic, sensory, and enteric neurons (Buj-Bello et al., Neuron 15, 821-828, 1995; Ebendal et al., J Neurosci Res 40, 276-284, 1995; Heuckeroth et al., Dev Biol 200, 116-29, 1998; Kotzbauer et al., Nature 384, 467-470, 1996; Trupp et al., J of Cell Biology 130, 137-148, 1995), persephin does not share any of these activities on peripheral neurons (Milbrandt et al., supra.

It was recently reported that GDNF and neurturin share receptors and signal transduction pathways (Creedon et al., *Proc. Natl. Acad. Sci. US 94*:7018-7023, 1997; Durbec et al., *Nature 381*:789-793, 1996; Trupp et al., *Nature 381*:785-789, 1996;

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Baloh et al., *Neuron 18*:793-802, 1997). These proteins act through a multicomponent receptor complex in which a transmembrane signal transducing component, the Ret protein-tyrosine kinase (Ret or Ret PTK), is activated upon the binding of a growth factor of the GDNF/neurturin family with a member of a family of closely related co-receptors named GFRα. A characteristic feature of the GFRα co-receptor family, is that its members have no transmembrane domain and are attached to the cell surface via a glycosyl-phosphatidylinositol (GPI) linkage (Durbec et al., *Nature 381*:789-793, 1996; Jing et al., *Cell 85*:1113-1124, 1996; Treanor et al., *Nature 382*:80-83, 1996; Trupp et al., *Nature 381*:785-789, 1996; Baloh et al., 1997, *supra*). The members of the GFRα family include GFRα1 (previously known as GDNFRα, TrnR1 and RetL1), GFRα2 (previously TrnR2, NTNRα and RetL2), GFRα3 (previously TrnR3) (GFRα Nomenclature Committee, *Neuron 19(3)*:485, 1997) and possibly GFRα4, a receptor currently only identified in the chicken (cGFRα4) (Enokido et al., *Current Biology 8*, 1019-1022, 1998).

15 Results from in vitro experiments by multiple groups together indicate that GFR $\alpha$ 1/RET is the preferred receptor for GDNF, and GFR $\alpha$ 2/RET is the preferred receptor for NTN, however cross-talk between the different receptors is possible (Baloh et al., 1997, supra; Jing et al., 1996, supra; Jing et al., J Biol. Chem. 272, 33111-33117, 1997; Klein et al., Nature 387, 717-721,1997; Sanicola et al., Proc. Natl. Acad. Sci., USA 94, 6238-6243,1997; Suvanto et al., Hum. Molec. Genet. 6, 20 1267-1273,1997; Treanor et al., 1996, supra). Recent analysis of GFRα1-deficient mice indicated that GFRa1 is the only physiologically relevant GDNF receptor in kidney organogenesis and enteric nervous system development (Cacalano et al., Neuron 21, 53-62,1998; Enomoto et al., Neuron 21, 317-324,1998). However, 25 GDNF-deficient mice have greater losses in peripheral ganglia than GFRα1-deficient mice, suggesting that GDNF can utilize additional other receptors to support survival of peripheral neurons, likely GFRα2/Ret (Cacalano et al., supra; Enomoto et al., supra). Persephin cannot signal through either the GFRα1/RET or GFRα2/RET receptor complexes (Milbrandt et al., supra), but a recent report indicated that 30 persephin binds to cGFRα4 and likely also signals through RET (Enokido et al., supra).

GFRα3 was first identified as an expressed sequence tag (EST) that is homologous to GFRa1 and GFRa2 (Baloh et al., Proc. Natl. Acad. Sci USA 95:5801-5806, 1998, supra; Jing et al., 1997, supra; Naveilhan et al., Proc Natl Acad Sci USA 95, 1295-300, 1998; Widenfalk et al., Eur. J. Neurosci. 10, 1508-1517, 1998; Worby et al., J Biol Chem 273, 3502-3508, 1998). However, analysis in transformed cells indicated that GFR \alpha 3 could not form a functional receptor with RET for any of the known GDNF family ligands, GDNF, neurturin or persephin (Baloh et al., 1998, supra;). GFR $\alpha$ 3 expression is much more restricted than GFR $\alpha$ 1 or GFR $\alpha$ 2, with high level expression observed only in developing peripheral nerve and ganglia (Baloh et al., 1998, supra; Naveilhan et al., supra; Widenfalk et al., supra; Worby et al., 1998). Furthermore, one report demonstrated that in sensory neurons of the trigeminal ganglion, GFRα3 is expressed in a population of neurons distinct from GFR $\alpha$ 1 and GFR $\alpha$ 2, and likely overlaps with RET (Naveilhan et al., *supra*). Although the structural similarity, expression, and functional data together suggested that GFR\alpha3 can interact with RET, the work reported herein was the first to demonstrate such an interaction and the first to identify a ligand for GFRa3.

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It is now generally believed that neurotrophic factors regulate many aspects of neuronal function, including survival and development in fetal life, and structural integrity and plasticity in adulthood. Since both acute nervous system injuries as well as chronic neurodegenerative diseases are characterized by structural damage and, possibly, by disease-induced apoptosis, it is likely that neurotrophic factors play some role in these afflictions. Indeed, a considerable body of evidence suggests that neurotrophic factors may be valuable therapeutic agents for treatment of these neurodegenerative conditions, which are perhaps the most socially and economically destructive diseases now afflicting our society. Nevertheless, because different neurotrophic factors can potentially act preferentially through different receptors and on different neuronal or non-neuronal cell types, there remains a continuing need for the identification of new members of neurotrophic factor families for use in the diagnosis and treatment of a variety of acute and chronic diseases of the nervous system.

### Summary of the Invention:

WO 00/18799

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Briefly, therefore, the present invention is directed to the new growth factor, artemin, which promotes cell survival and growth. Like GDNF and neurturin, artemin supports the survival of several peripheral neuron populations, and can also support dopaminergic neurons of the ventral midbrain. However, artemin is the only member of the GDNF family that binds to GFRα and activates the GFRα3/RET receptor complex and, in addition, like GDNF and neurturin, artemin also binds to and activates GFRα1/RET. GFRα3 is described in patent application entitled "GFRα3, A Novel Member of the GDNF Coreceptor Family" filed concurrently with the present application on December 22, 1998 and this GFRα3 application is incorporated herein in its entirety by reference.

The present invention thus provides artemin polypeptides and polynucleotides. Artemin polypeptides within the scope of the present invention include any of the naturally occurring artemin polypeptides, conservatively substituted variants thereof or fragments thereof. The artemin polypeptides promote survival of peripheral and central neurons in culture, preferably in any one of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosine-hydroxylase-expressing dopaminergic ventral midbrain neurons; more preferably, in any combination of these neurons and, most preferably, in all of these neurons.

Preferred artemin polypeptides include the predicted human mature polypeptides (SEQ ID NOS:3-5, see Figs. 3A, 3B, and 3C), and the predicted mouse mature polypeptides (SEQ ID NOS:34-36). These mature polypeptides can be generated by cleaving human or mouse pre-pro artemin (SEQ ID NOS: 26 and 29, respectively) or human or mouse pro-artemin (SEQ ID NOS:40 and 41, respectively), which are also within the scope of this invention, at one of the RXXR cleavage sites (see Figs. 1A, 1B, 1C and 2B). In addition, mature artemin can be generated by cleaving a polypeptide containing an N-terminal non-artemin pre-pro- region a mature artemin. Artemin polypeptides also include fragments from the first canonical cysteine to the seventh canonical cysteine of human or mouse mature artemin (SEQ ID NO:19 and SEQ ID NO:33, respectively).

Artemin is believed to show at least 75% sequence identity among homologous sequences from different mammalian species and, hence, mammalian

WO 00/18799 PCT/US99/22604

orthologs of mature artemin are believed to have at least 75% sequence identity with mature human artemin (SEQ ID NOS:3-5) and mature mouse artemin (SEQ ID NOS:34-36). Sequence homology may be as low as 65% in non-mammalian species such as avian species.

Human artemin polypeptide is encoded by the cDNA contained in the clone deposited with the ATCC on December 22, 1998. Thus, human artemin polypeptide can be produced by transforming a host cell with the cDNA of this clone operably linked to expression regulatory elements. Conditions are then provided such that the cell expresses the encoded human artemin polypeptide. Such human artemin polypeptides are within the scope of the present invention.

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The present invention also provides compositions comprising an artemin polypeptide and a pharmaceutically acceptable carrier suitable for administering to a cell to provide trophic support and/or to produce differentiation of that cell *in vitro* or  $ex\ vivo$  or to a cell in a patient. In order to facilitate the growth promoting effect of artemin, a GFR $\alpha$  coreceptor can be administered with artemin. Thus, in another embodiment, the present invention provides compositions which comprise an artemin polypeptide and a GFR $\alpha$  polypeptide such as GFR $\alpha$ 3 or GFR $\alpha$ 1.

In another embodiment, the present invention includes pan-growth factors and polynucleotides encoding the pan-growth factors. The pan-growth factors comprise a portion of the artemin polypeptide and a portion of at least one other growth factor from the TGF- $\beta$  superfamily. Preferably, the other growth factor can be neurturin, persephin or GDNF.

The present invention in another embodiment also provides isolated and purified and/or recombinant nucleic acid molecules comprising artemin nucleotide sequences. The artemin nucleotide sequences encode polypeptides consisting of artemin polypeptides, conservatively substituted variants thereof and fragments thereof. Also within the scope of this invention are fragments of artemin nucleotide sequences of at least 15 contiguous nucleotides. The artemin nucleic acid molecules specifically hybridize to mature human artemin nucleotide sequences (SEQ ID NOS:6-8) or their complements (SEQ ID NOS:9-11) or to mature mouse artemin nucleotide sequences (SEQ ID NOS:60-62).

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Preferred polynucleotides of the present invention comprise the human polynucleotides which encode mature human artemin polypeptides (SEQ ID NOS:3-5), in particular, the polynucleotide sequences in SEQ ID NOS: 6-8 as depicted in Figs. 3A, 3B, and 3C or the corresponding mouse polynucleotides which encode mature mouse artemin polypeptides. (SEQ ID NOS:34-36), in particular, the polynucleotide sequences in SEQ ID NOS:37-39. Also encompassed by the present invention are human and mouse pro-artemin polynucleotides which encode proartemin polypeptides (SEQ ID NOS:40 and 41, respectively), in particular, the polynucleotide sequences in SEQ ID NOS:42 and 43, respectively, and human and mouse pre-pro artemin polynucleotides which encode pre-pro artemin polypeptides (SEO ID NOS:26 and 29, respectively), in particular, the human polynucleotide sequences, SEQ ID NOS:24, SEQ ID NO:30 or SEQ ID NO:46, and the mouse polynucleotide sequence, SEQ ID NO:27. In addition, polynucleotides can encode a polypeptide containing an N-terminal non-artemin pre-pro- region and a mature artemin such that upon expression of the polypeptide in an expression system and cleavage of the N-terminal non-artemin pre-pro region, a mature artemin is produced. Artemin polynucleotides also include portions of the artemin polynucleotide sequence encoding the polypeptide from the first canonical cysteine to the seventh canonical cysteine (SEQ ID NOS:19 and 33).

Sequences that are capable of hybridizing to human or mouse mature artemin polynucleotides can be used in methods for detecting the artemin gene and transcription products thereof, as well as in isolating artemin-encoding polynucleotides from other mammalian and non-mammalian species. Such methods are also within the scope of the present invention.

The present invention also provides antibodies which specifically react with artemin or a fragment thereof and methods for detecting artemin polypeptide in a sample by binding to the antibody.

In another embodiment, the present invention also provides methods for providing trophic support to and/or for producing differentiation of a target cell which comprises treating the cell with an effective amount of an artemin polypeptide. The target cell can be given artemin *in vitro*, *ex vivo* or *in vivo* in a patient suffering from a medical condition which causes degeneration or a loss of normal function of the target

WO 00/18799

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cell. In one preferred embodiment, the cell is treated *in vivo* by administering to the patient an artemin polypeptide or an artemin polynucleotide encoding for expression the artemin polypeptide. The patient can be suffering from disease such as peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, nervous system tumors such as neuroblastomas, multiple sclerosis, infection, small cell lung carcinoma and enteric diseases such as idiopathic chronic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers.

Among the several advantages found to be achieved by the present invention, therefore, may be noted the provision of a new growth factor, artemin, in particular human artemin, which can be used to prevent the atrophy, degeneration or death of certain cells, in particular, neurons in need of trophic support; the provision of polynucleotides encoding artemin for use in gene therapy; the provision of methods for obtaining artemin by recombinant techniques; the provision of methods for providing trophic support to target cells, particularly neurons; the provision of methods for treating disease conditions involving cellular degeneration, and in particular, neuronal degeneration; the provision of methods that can detect and monitor artemin levels in a patient; and the provision of methods that can detect alterations in the artemin gene.

### Brief Description of the Drawings

Figure 1A illustrates complementary human artemin genomic nucleotide sequences (SEQ ID NOS:1-2) which were assembled as described in the text, with the amino acids (SEQ ID NO:12) encoded by one of the reading frames shown below the nucleotide sequences and the possible RXXR proteolytic processing sites indicated by boxes.

Figure 1B illustrates the complementary cDNA sequences (SEQ ID NOS:24 and 25) and amino acid sequence (SEQ ID NO:26) of human pre-pro-artemin.

Figure 1C illustrates the complementary cDNA sequences (SEQ ID NOS:27 and 28) and amino acid sequence (SEQ ID NO:29) of mouse pre-pro-artemin.

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Figure 1D illustrates the complementary cDNA sequences (SEQ ID NOS:30 and 31) and amino acid sequence (SEQ ID NO:32) starting from a second methionine found in some human artemin cDNA clones which represent alternatively spliced human artemin mRNA;

Figure 2A illustrates an alignment of the amino acid sequences for the predicted mature forms of human GDNF (hGDNF, SEQ ID NO:13), human neurturin (hNTN, SEQ ID NO:14), human persephin (hPSP, SEQ ID NO:15) and human artemin (hART, SEQ ID NO:3) with identical amino acid residues enclosed in boxes and gaps inserted by the alignment program indicated by dashes;

Figure 2B illustrates the alignment of human and mouse pre-pro artemin (SEQ ID NOS:26 and 29, respectively) with the human and mouse putative signal sequence, pre- region, shaded (amino acids 1-39, SEQ ID NO:51 and SEQ ID NO:52, respectively), putative RXXR cleavage sites designated by thick lines, and arrowheads denoting the location of the two introns in the artemin gene.

Figure 2C is a schematic diagram of the location of the two introns in the artemin gene, and their splicing to produce artemin mRNA, with the location of a second starting methionine identified by RACE PCR in cDNA's isolated from some human cDNA libraries indicated by Met\*.

Figure 3A illustrates the amino acid sequence (SEQ ID NO:3) and the coding sequence (SEQ ID NO:6) of a first predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:9).

Figure 3B illustrates the amino acid sequence (SEQ ID NO:4) and the coding sequence (SEQ ID NO:7) of a second predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:10).

Figure 3C illustrates the amino acid sequence (SEQ ID NO:5) and the coding sequence (SEQ ID NO:8) of a third predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:11).

Figure 4 illustrates the family member sequence identity in the region between the first and seventh canonical framework cysteine residues aligned beginning with the first canonical framework cysteine for human GDNF (hGDNF, SEQ ID NO:16), human neurturin (hNTN, SEQ ID NO:17), human persephin (hPSP, SEQ ID NO:18) and human artemin (hART, SEQ ID NO:19).

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Figure 5 illustrates expression of artemin in human adult and fetal tissues showing in Fig. 5A an RNA spot blot containing poly(A) RNA's isolated from the human tissues shown in Fig. 5B and probed with a fragment of the human artemin cDNA.

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PCT/US99/22604

Figures 6A and 6B illustrate *in situ* hybridization analysis of artemin expression in embryonic rat using a <sup>32</sup>P-labeled RNA probe generated from a rat artemin cDNA fragment as a probe, showing in Fig. 6A a sagittal section of an embryonic day 14 (E14) rat, with the section oriented such that the top is the rostral and the left side is the dorsal, in which no signal is observed in the dorsal root ganglia (DRG) and strong signal is observed in probable exiting nerve roots (NR), with the arrowhead indicating expression observed in tissue below the liver that likely represents a lateral extension of the superior mesenteric artery, and showing in Fig. 6B a parasagittal section of E14 rat in the same orientation as in (A), in which high level artemin expression surrounding the developing superior mesenteric artery (SMA) is detected.

Figure 6C illustrates semi-quantitative RT-PCR analysis of artemin expression in Schwann cells isolated from neonatal rats (PS), from the adult sciatic nerve (N), and from the distal segment of the sciatic nerve at the indicated times following nerve transection.

Figure 7 illustrates that artemin supports the survival of peripheral and central neurons in culture, in which: Figs 7A and 7B show histograms of the number of surviving dorsal root ganglion (DRG) neurons cultured from post-natal day 1 (P1) rats in the presence of 50 ng/ml of the indicated factors (Fig. 7A, mean and SEM are shown, n=5-11) or in the presence of the indicated amounts of artemin (Fig. 7B, mean and SEM are shown, n=5); Fig. 7C shows a histogram of number of surviving trigeminal ganglion (TG) neurons cultured from P1 rats in the presence of 50 ng/ml of the indicated growth factor (mean and SEM are shown, n=4); Fig. 7D shows a histogram of the number of surviving nodose ganglion (NG) neurons cultured from P0 rats in the presence of BDNF (100 ng/ml) or the indicated GDNF family ligands (50 ng/ml) (mean and SEM from a representative experiment are shown, n=2); Fig. 7E shows a histogram of the number of surviving superior cervical ganglion (SCG) from P0 rats maintained in the presence of NGF for 5 days and then cultured in the

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presence of 50 ng/ml of the indicated factors (mean and SEM are shown, n=2-8); and Fig. 7F shows a histogram of the number of tyrosine hydroxylase-expressing (TH+) dopaminergic ventral midbrain neurons from E14 rats (EVM) cultured in the presence of 50 ng/ml of the indicated factors (mean and SEM are shown, n=9-15).

Figure 8 illustrates that artemin induces differentiation and proliferation in neuroblastoma cell lines showing in Figs. 8A-8C photographs of SK-SY5Y neuroblastoma cells cultured in the presence of no factor (Cn) (Fig. 8A), 10 μM all-trans retinoic acid (RA), a known inducer of differentiation of these cells (Fig. 8B), or 50 ng/mL artemin (ART) (Fig. 8C), and in Fig. 8D a histogram of BrdU incorporation by NBL-S neuroblastoma cells in the presence of 50 ng/mL of the indicated factor.

Figure 9A illustrates that artemin activates RET and downstream signaling in primary cultured SCG neurons showing an immunoblot of tyrosine phosphorylated RET or phosphorylated MAPK (MAPK) in lysates from SCG neurons treated with either no factor (Cn), GDNF, or artemin (ART) at 50 ng/ml.

Figure 9B illustrates that artemin activates RET in NBL-S neuroblastoma cells, showing an immunoblot of tyrosine phosphorylated RET or phosphorylated MAPK (MAPK) in lysates from NBL-S cells stimulated as in Fig. 9A in the absence or presence of the enzyme PI-PLC, which specifically cleaves GPI-anchored proteins from the cell surface.

Figures 9C-9E illustrate the direct binding of GFRα1-Fc, GFRα2-Fc, and GFRα3-Fc receptor bodies to GDNF ligand family members showing graphs of the amount of absorbance at 450 nm plotted against increasing concentrations of the indicated soluble GFRα-Fc fusion protein added to microtiter plates coated with GDNF, neurturin, artemin or persephin, which were then treated with anti-human Fc antibodies conjugated to horseradish peroxidase (HRP) and binding of receptor bodies measured using the chromogenic HRP substrate 3,3',5,5'-tetramethylbenzidine, with error bars representing the standard deviation of duplicates from a representative experiment.

Figure 10 illustrates receptor activation by GDNF ligand family members in the presence of defined coreceptor components showing bar graphs of the amount of luciferase expression induced by the indicated growth factor in RET-expressing MG87 fibroblasts (RET-3T3) (Fig. 10A) or in NLF neuroblastoma cells (NLF) (Fig.

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10B) transiently transformed with the Gal4-Elk/Gal4-Luc reporter system together with an expression plasmid for the indicated coreceptor or the CMV plasmid with no insert (Control), with fold-activation determined by dividing luciferase activity in the indicated treatment condition by the no treatment control, and error bars representing the standard deviation of duplicate measurements from a representative experiment.

Figure 11 is a schematic diagram of ligand/receptor interactions in the GDNF ligand family deduced from multiple experimental paradigms *in vitro*, with large arrows indicating preferred ligand receptor interactions and smaller arrows indicating alternate receptor interactions.

Figure 12 illustrates the alignment of the amino acid sequences of human and mouse GFR $\alpha$ 3 precursor (SEQ ID NOS:63 and 64, respectively), with identical residues enclosed in boxes, the N-terminal signal sequence shaded (amino acids 1-31 in hGFR $\alpha$ 3 and 1-28 in mGFR $\alpha$ 3), mature GFR $\alpha$ 3 (amino acids 32-372 in hGFR $\alpha$ 3 and 29-369 in mGFR $\alpha$ 3), a shaded C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide shaded (amino acids 373-400 of hGFR $\alpha$ 3 and 370-397 of mGFR $\alpha$ 3) and putative N-linked glycosylation sites marked by black dots.

Figure 13 illustrates the human nucleotide sequence encoding GFR $\alpha$ 3 precursor (SEQ ID NO:65) which includes the encoding sequence for the N-terminal signal sequence (nucleotides 1-93), the encoding sequence for mature GFR $\alpha$ 3 (nucleotides 94-1116) and the encoding sequence for the C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide (nucleotides 1117-1203).

Figure 14A-C illustrate the complementary human genomic sequence that includes the artemin gene (SEQ ID NOS: 68 and 69) and the sequences of the three reading frames (SEQ ID NOS:70-72).

Figure 15 illustrates the partial rat artemin cDNA complementary sequences (SEQ ID NOS:73 and 74) and the encoded amino acid sequence (SEQ ID NO:75).

#### Description of the Preferred Embodiments

The present invention is based upon the identification, isolation and sequencing of genomic and cDNA clones that encode a new member of the neurturin/persephin/ GDNF ligand family identified herein as artemin (referenced in copending application 09/163,283 as GF4). The discovery of artemin follows that of

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neurturin which is described in copending U.S. Application Serial No. 08/775,414, which is incorporated in its entirety by reference, and persephin which is described in copending U.S. Applications Serial Nos. 08/981,739 and 08/931,858 both of which are incorporated in their entirety by reference.

14

Artemin promotes the survival of several peripheral and central neuronal populations *in vitro*, including sympathetic neurons, neural crest, placodally-derived sensory neurons, and dopaminergic midbrain neurons. Artemin activates both the  $GFR\alpha 3/RET$  and  $GFR\alpha 1/RET$  receptor complexes *in vitro*, but it is believed  $GFR\alpha 3/RET$  is the preferred multicomponent receptor for artemin signaling *in vivo*.

As described in more detail below, artemin was discovered by using the murine pre-pro neurturin amino acid sequence to scan the High Throughput Genome Sequences (hgts) database. This search identified two hgts sequences (AC005038 and AC005051) which had regions of DNA homologous to a DNA encoding pre-pro neurturin. Although, the hgts sequences were later discovered to have numerous sequence errors, i.e. omissions, additions and incorrect bases, one of the hgts sequences (AC005038) had a stretch of 197 nucleotides (nucleotides 67,464 to 67,660) which ultimately turned out to be identical to nucleotides 663-467 of the complementary strand of the Artemin nucleic acid (Fig. 1B) and the other (AC005051) had a stretch of 183 nucleotides (nucleotides 113,379 to 113,561) identical to nucleotides 648-468 of the complementary strand of the Artemin nucleic acid (Fig. 1B).

In order to determine whether the hgts sequences corresponded to coding regions of a new growth factor, oligonucleotide primers were designed based upon the hgts sequences to amplify overlapping polynucleotides of the suspected coding region from human genomic DNA. The overlapping polynucleotides were sequenced and this sequence information was combined with the hgts sequences to assemble the complementary genomic nucleotide sequences (SEQ ID NO:1-2) shown in Fig. 1A. One of the reading frames in SEQ ID NO:1 encodes a partial pro-artemin amino acid sequence that extends from within a pro-domain including three potential RXXR proteolytic processing sites (boxes in Fig. 1A) and terminates at the second glycine residue shown in the last line of Figure 1A. Alignment of the predicted mature human artemin (hART) protein with the other GDNF ligands (Figure 2A) confirmed that

artemin represents a new member of the GDNF ligand family, with artemin being most similar to neurturin and persephin (~45% identity), and somewhat more divergent from GDNF (~36% identity). Primers designed from the human artemin genomic sequence were used to identify a clone containing the mouse artemin gene.

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To isolate full length mRNA species encoding artemin, primers corresponding to the human and mouse artemin genomic sequences were used to perform rapid amplification of cDNA ends (RACE) PCR on multiple human and mouse tissue cDNA libraries. Analysis of the full-length cDNAs from human and mouse (Figs 1B and 1C) enabled the prediction of full-length human and mouse artemin proteins (Figs. 1B, 1C and 2B) having a signal peptide for secretion (amino acids 1- 39 of Fig. 2B), and a large pro-region separated from the mature region by multiple conserved RXXR furin protease cleavage sites (Fig. 2B). Comparison of the human mRNA sequence with the genomic locus revealed the presence of two introns in the artemin coding region (Figure 2B and 2C), the second of which is positioned similarly to the introns found in the pro-domains of other GDNF family ligands.

Reference to mature artemin herein is intended to be construed to include growth factors of any origin which are substantially homologous to and which are biologically equivalent to mature human and mouse artemin characterized and described herein. Such substantially homologous growth factors may be native to any tissue or species and, similarly, biological activity can be characterized in any of a number of biological assay systems. Reference to pre-pro-artemin is intended to be construed to include pre-pro growth factors containing a pre- or leader or signal sequence region, a pro- sequence region and a mature artemin amino acid sequence as defined herein. Reference to pro-artemin is intended to mean a polypeptide lacking the signal sequence region, but containing both a pro-region ending in an RXXR cleavage sequence and a mature artemin amino acid sequence.

The terms "biologically equivalent" are intended to mean that the compositions of the present invention are capable of demonstrating some or all of the same growth promoting properties in a similar fashion, although not necessarily to the same degree as the recombinantly produced human artemin identified herein.

By "substantially homologous" it is meant that the degree of sequence identity between artemin orthologs including human, mouse and artemin from any other

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species, is greater than that between paralogs such as human artemin and human neurturin or human artemin and human persephin, and greater than that reported previously for members of the TGF-ß superfamily (for discussion of homology of TGF-ß superfamily members see Kingsley, *Genes and Dev* 8:133-46, 1994).

Sequence identity or percent identity is intended to mean the percentage of same residues between two sequences aligned using the Clustal method (Higgins et al, Cabios 8:189-191, 1992) of multiple sequence alignment in the Lasergene biocomputing software (DNASTAR, INC, Madison, WI). In this method, multiple alignments are carried out in a progressive manner, in which larger and larger alignment groups are assembled using similarity scores calculated from a series of pairwise alignments. Optimal sequence alignments are obtained by finding the maximum alignment score, which is the average of all scores between the separate residues in the alignment, determined from a residue weight table representing the probability of a given amino acid change occurring in two related proteins over a given evolutionary interval. Penalties for opening and lengthening gaps in the alignment contribute to the score. The default parameters used with this program are as follows: gap penalty for multiple alignment = 10; gap length penalty for multiple alignment = 10; k-tuple value in pairwise alignment = 1; gap penalty in pairwise alignment = 3; window value in pairwise alignment = 5; diagonals saved in pairwise alignment = 5. The residue weight table used for the alignment program is PAM250 (Dayhoff et al., in Atlas of Protein Sequence and Structure, Dayhoff, Ed., NBRF, Washington, Vol. 5, suppl. 3, p. 345, 1978).

To determine percent sequence identity between two sequences, the number of identical amino acids in the aligned sequences is divided by the total number of amino acids in the reference sequence. As used herein, the reference sequence is human artemin when determining its percent identity with mouse artemin and with the non-artemin growth factors, human neurturin, human GDNF or human persephin. Similarly, the reference sequence would be mouse artemin when determining its percent identity with mouse GDNF, mouse neurturin or mouse persephin and the reference sequence would be rat artemin when determining its percent identity with rat GDNF, rat neurturin and rat persephin. Referencing is to human neurturin when

17

PCT/US99/22604

determining percent identity between human neurturin and non-human neurturin or between human neurturin and its human paralogs GDNF and persephin.

Percent conservation is calculated from the above alignment by adding the number of identical residues to the number of positions at which the two residues represent a conservative substitution (defined as having a log odds value of greater than or equal to 0.3 in the PAM250 residue weight table) and dividing by the total number of amino acids in the reference sequence. Preferred conservative amino acid changes are: R-K; E-D, Y-F, L-M; V-I, Q-H.

Table 1 shows the percent identity (I) and conservation (C) for comparisons of mature artemin, mature persephin, mature neurturin and mature GDNF from various species. Comparisons were made between mature human artemin (hART) and mature mouse artemin (mART) and between mature human artemin and mature human persephin (hPSP), mature human neurturin (hNTN) or mature human GDNF (hGDNF) using the alignments shown in Figs. 2A and 2B and the preferred conservative substitutions stated above. The persephin and neurturin comparisons were made as described in copending application Serial No. 08/931,858, with the first listed sequence being the reference sequence.

PCT/US99/22604

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Table 1.

	COMPARISON	% IDENTITY	% CONSERVATION	
	hART v. mART	88	90	
	hART v. hPSP	45	48	
5	hART v. hNTN	49	51	
	hART v. hGDNF	36	40	
	hPSP v. mPSP	81	81	
	hPSP v. rPSP	80	81	
10	mPSP v. rPSP	94	96	
	hPSP v. hNTN	49	50	
	hPSP v. hGDNF	40	43	
	hNTN v. mNTN	90	93	
15	hNTN v. rGDNF	44	53	
	hNTN v. mGDNF	43	52	
	hNTN v. hGDNF	43	53	
	mNTN v. rGDNF	42	52	
	mNTN v. mGDNF	41	51	
20	mNTN v. hGDNF	41	52	

The sequence identity between human artemin and mouse artemin is about 88%. The persephin comparisons shown in Table 1 indicate that the identity between human persephin and mouse or rat persephin is about 80% whereas the degree of identity between mouse and rat persephin is about 94%. The neurturin comparisons in Table 1 indicate that mature mouse and human neurturin proteins have about 90% sequence identity. Furthermore, all artemin, persephin and neurturin orthologs of non-human mammalian species are believed to similarly have at least about 75% sequence identity with human artemin, human persephin, or human neurturin, respectively. For artemin, persephin or neurturin orthologs from non-mammalian species such as avian species, it is believed that the degree of homology with human artemin, human persephin, or human neurturin is at least about 65% identity.

WO 00/18799 PCT/US99/22604

19

By way of comparison, the variations between family members of the GDNF ligand family of growth factors can be seen by the comparisons shown in Table 1. For example, human artemin has about 49% sequence identity with human neurturin and about 36% sequence identity with human GDNF. Human persephin has about 49% sequence identity with human neurturin and about 40% sequence identity with human GDNF. Similarly, human neurturin has about 43% sequence identity with human GDNF. Thus, it is believed that any other member of the GDNF ligand family will have a similar sequence identity of about 40% of that of artemin, neurturin, persephin or GDNF of the same species and within a range of about 30% to about 60% sequence identity with artemin, neurturin, persephin or GDNF of the same species.

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Based on the data in Table 1, a given member of the GDNF ligand family would be expected to have lesser sequence identity with any other family member of the same species than the sequence identity present in orthologs of that family member in other species just as human GDNF and human neurturin are more closely related to mouse GDNF and mouse neurturin, respectively, than to each other or to GDNF (see U.S. Patent NO. 5,739,307). Similarly, any given member of the GDNF ligand family would be expected to have greater sequence identity with another family member than to any other known member of the TGF- $\beta$  superfamily (Kingsley, *supra*).

The conservation between members of the GDNF ligand family is also seen in Figure 4, which shows an alignment, using the Clustal program described above, of the human sequences for GDNF, neurturin (NTN), persephin (PSP) and artemin (ART) from the first to the seventh framework cysteine residues. From this alignment, it is evident that artemin is closely related to the other family members in that artemin and persephin share 46 out of 96 residues (48%), artemin shares 48 out of 96 residues (50%) with neurturin, and artemin and GDNF share 36 out of 96 residues (38%).

Orthologs of pre-pro artemin in non-human mammalian species can be identified by virtue of the mature portion of the amino acid sequence having at least about 75% sequence identity with one of the predicted mature human artemin sequences disclosed herein and nonmammalian orthologs of pre-pro artemin can be

identified by virtue of the mature portion of the amino acid sequence having at least about 65% identity with a mature human artemin sequence.

Also included within the meaning of substantially homologous is any artemin polypeptide which may be isolated by virtue of cross-reactivity with antibodies specific to human artemin or whose encoding nucleotide sequences, including genomic DNA, mRNA or cDNA may be isolated through hybridization with the complementary sequences shown in Figs 1-3 or fragments thereof. It will also be appreciated by one skilled in the art that degenerate DNA sequences can encode human artemin and these are also intended to be included within the present invention.

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Conservatively substituted artemin proteins retaining the biological activity of naturally occurring artemin are also within the scope of the present invention. Conservative amino acid substitutions refer to the interchangeability of residues having similar side chains. Conservatively substituted amino acids can be grouped according to the chemical properties of their side chains. For example, one grouping of amino acids includes those amino acids have neutral and hydrophobic side chains (A, V, L, I, P, W, F, and M); another grouping is those amino acids having neutral and polar side chains (G, S, T, Y, C, N, and Q); another grouping is those amino acids having basic side chains (K, R, and H); another grouping is those amino acids having acidic side chains (D and E); another grouping is those amino acids having aliphatic side chains (G, A, V, L, and I); another grouping is those amino acids having aliphatic-hydroxyl side chains (S and T); another grouping is those amino acids having amine-containing side chains (N, Q, K, R, and H); another grouping is those amino acids having aromatic side chains (F, Y, and W); and another grouping is those amino acids having sulfur-containing side chains (C and M). Preferred conservative amino acid substitutions groups are: R-K; E-D, Y-F, L-M; V-I, and Q-H.

As used herein, an artemin polypeptide can also include modifications of the artemin sequences identified herein, including sequences in which one or more amino acids have been inserted, deleted or replaced with a different amino acid or a modified or unusual amino acid, as well as modifications such as glycosylation or phosphorylation of one or more amino acids so long as the polypeptide containing the modified sequence retains the biological activity of artemin. Inserted or deleted

WO 00/18799 PCT/US99/22604

21

amino acid(s) can be added to or removed from the N-terminus, C-terminus or within the naturally-occurring amino acid sequence. By retaining the biological activity, it is meant that the modified polypeptide can bind to and activate GFRα1/RET and/or GFRα3/RET expressed by a cell, although not necessarily at the same level of potency as that of the mature human artemin polypeptide identified herein. Assays for testing such activation are known in the art and include the Gal4-Elk/Gal4-Luc reporter system described in Example 7 below. The term "trophic support" is used herein to mean that a growth factor such as artemin provides nourishment to a cell such that the cell maintains or recovers at least one of its normal functions.

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It is intended that the term artemin polypeptide also includes naturally occurring allelic variants of the human artemin sequences disclosed herein. For example, cDNAs encoding a second starting methionine were identified by RACE PCR from some human cDNA libraries and would encode the pre-pro-artemin shown in Fig. 1D (SEQ ID NO:32). However, mature artemin generated from this variant would be identical to the mature human artemin shown in Fig. 3A (SEQ ID NO:3).

Fragments of artemin are also encompassed by the present invention. Such fragments may be of any length but preferably retain the biological activity of artemin or are antigenic. The minimum length of such biologically active or antigenic fragments can readily be determined by those skilled in the art using known techniques. Preferably, the minimum length of fragments of artemin is at least 8 amino acids, more preferably, at least 10 amino acids, still more preferably at least 12 amino acids, even still more preferably at least 15 amino acids and most preferably, at least 20 amino acids or greater. One fragment of artemin which is believed to retain the survival promoting activity of artemin begins at the first conserved cysteine residue and ends at the seventh conserved cysteine residue (Fig. 4, SEO ID NO:19 (human) or SEQ ID NO:33 (mouse)). Antigenic fragments are capable of eliciting artemin-specific antibodies when administered to a host animal and includes those smaller fragments that require conjugation to a carrier molecule to be immunogenic. Typically, antigenic fragments will be at least 5 or 6 amino acids in length and may be any length up to the length of mature artemin, preferably an antigenic fragment is 8, more preferably 10, still more preferably 12 amino acids in length, even still more

WO 00/18799 PCT/US99/22604

22

preferably 15 amino acids in length and most preferably 20 amino acids or more in length.

It is also believed that particular discrete fragments of artemin, or analogues thereof, can serve as an agonist where the fragment activates an artemin receptor, such as  $GFR\alpha 3/RET$  or  $GFR\alpha 1/RET$ , to elicit the survival or growth promoting action on a target cell, or other artemin fragments or analogues can serve as antagonists to artemin where they bind to, but do not activate, the receptor or do not promote survival and growth. Such fragments or analogues that are agonists and those that are antagonists are also within the scope of the present invention.

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Although it is not intended that the inventors herein be bound by any theory, it is thought that the human artemin protein identified herein as well as orthologs from other tissues and species may exist as dimers in their biologically active form in a manner consistent with what is known for other factors of the TGF-ß superfamily.

In addition to homodimers, the monomeric units of artemin dimers can be used to construct stable growth factor heterodimers or heteromultimers comprising at least one monomer unit of artemin. This can be done by dissociating a homodimer of artemin into its component monomeric units and reassociating in the presence of a monomeric unit of a second or subsequent homodimeric growth factor. This second or subsequent homodimeric growth factor can be selected from a variety of growth factors including neurturin, GDNF, persephin, a member of the NGF family such as NGF, BDNF, NT-3 and NT-4/5, a member of the TGF-β superfamily, a vascular endothelial growth factor, a member of the CNTF/LIF family and the like. By forming heterodimers or heteromultimers of artemin and one or more other growth factors, the resultant hybrid growth factor would be expected to be able to bind to at least two distinct receptor types preferentially having a different tissue distribution. The resultant heterodimers or heteromultimers would be expected to show a different and, possibly, an enlarged spectrum of cells upon which it could act or to provide greater potency. It is also possible that the heterodimer or heteromultimer might provide synergistic effects not seen with homodimers or homomultimers. For example, the combination of factors from different classes has been shown to promote long-term survival of oligodendrocytes whereas single factors or combinations of

factors within the same class promoted short-term survival (Barres et al., *Development 118*:283-295, 1993).

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Heterodimers can be formed by a number of methods. For example, homodimers can be mixed and subjected to conditions in which dissociation/unfolding occurs, such as in the presence of a dissociation/unfolding agent, followed by subjection to conditions which allow monomer reassociation and formation of heterodimers. Dissociation/unfolding agents include any agent known to promote the dissociation of proteins. Such agents include, but are not limited to, guanidine hydrochloride, urea, potassium thiocyanate, pH lowering agents such as buffered HCl solutions, and polar, water miscible organic solvents such as acetonitrile or alcohols such as propanol or isopropanol. In addition, for homodimers linked covalently by disulfide bonds as is the case with TGF-β family members, reducing agents such as dithiothreitol and β-mercaptoethanol can be used for dissociation/unfolding and for reassociation/refolding.

Heterodimers can also be made by transforming a cell with two or more factors such that the transformed cell produces heterodimers as has been done with the neurotrophins. (Heymach and Schooter, *J Biol Chem 270*:12297-12304, 1995). Another method of forming heterodimers is by combining artemin homodimers and a homodimer from a second growth factor and incubating the mixture at 37°C.

When heterodimers are produced from homodimers, the heterodimers may then be separated from homodimers using methods available to those skilled in the art such as, for example, by elution from preparative, non-denaturing polyacrylamide gels. Alternatively, heterodimers may be purified using high pressure cation exchange chromatography such as with a Mono S cation exchange column or by sequential immunoaffinity columns.

It is well known in the art that many proteins are synthesized within a cell with a signal sequence at the N-terminus of the mature protein sequence and the protein carrying such a leader sequence is referred to as a preprotein. The pre- portion of the protein is cleaved during cellular processing of the protein. In addition to a pre-leader sequence, many proteins contain a distinct pro sequence that describes a region on a protein that is a stable precursor of the mature protein. Proteins synthesized with both pre- and pro- regions are referred to as preproproteins. In view of the processing

WO 00/18799 PCT/US99/22604

24

events known to occur with other TGF-ß family members, the inventors believe that the form of the artemin protein as synthesized within a cell is the pre-pro-artemin.

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Human and mouse pre-pro-artemin polypeptides are believed to preferably contain an N-terminal methionine of a signal sequence (pre-region) of 39 amino acids (Fig. 2B; SEQ ID NOS:48 and 49, respectively; amino acids 1-39 of SEO ID NOS:26 and 29, respectively). It is known that the full length of a leader sequence is not necessarily required for the sequence to act as a signal sequence and, therefore, within the definition of pre-region of artemin is included fragments thereof, usually Nterminal fragments, that retain the property of being able to act as a signal sequence, that is to facilitate co-translational insertion into the membrane of one or more cellular organelles such as endoplasmic reticulum, mitochondria, golgi, plasma membrane and the like. It is also possible that the pre-pro-artemin polypeptide might have one or more isoforms having an N-terminal Leucine, inasmuch as isoforms of another growth factor, human fibroblast growth factor-2, have alternative initiations of translation at CUG start codons in addition to AUG (Arnaud et al., Molecular and Cellular Biology 19:505-514, 1999). Thus, isoforms of pre-pro-artemin could begin. for example, before the methionine (the first amino acid shown in Figure 1B) with the start point being at the CTG codon beginning at nucleotide 284 in the genomic human artemin sequence (see Figure 14) or after the same methionine with the start point being the CTG codon beginning at nucleotide 329 so long as the pre-region of the isoform can serve as a signal sequence. All such isoforms are within the scope of the present invention.

Moreover, also within the scope of the present invention are polypeptides containing a non-artemin pre-pro- region and a mature artemin as well as polynucleotides encoding such polypeptides. The polypeptides can generate a mature artemin upon cleavage of the non-artemin pre-pro- region and the polynucleotides can be used in an expression system to produce the polypeptide which upon cleavage of the non-artemin pre-pro- region yields mature artemin. Such non-artemin pre-pro- region polypeptides and encoding polynucleotides are well known in the art and, preferably, include a pre-pro- region from another growth factor such as neurturin or persephin although any pre-pro- region can be used so long as it functions as a pre-pro- region as described herein.

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The artemin pre- region is followed by a pro- domain which is believed to preferably terminate with an RXXR consensus site immediately before the N-terminal amino acid of mature artemin. Thus, mature human and mouse artemin are believed to be preferably generated by proteolytic cleavage of human and mouse pro-artemin (SEQ ID NOS:40 and 41, respectively) after one of the three RXXR consensus sequences shown in Figs. 1A and 2B, (RARR, RGGR and RAAR for human and RTLR, RGAR and RAAR for mouse) although it is possible that cleavage could occur at some other, non-consensus site to produce additional isoforms. All such isoforms are included within the scope of the present invention.

Cleavage after the first RXXR sequence following the N-terminus of human pro-artemin will generate the mature polypeptide shown in Fig. 3C (SEQ ID NO:5). Similarly, cleavage after the second and third RXXR sequences will generate the mature human polypeptides shown in Figs. 3B (SEQ ID NO:4) and 3A (SEQ ID NO:3), respectively. By analogy, predicted mature mouse artemin sequences are SEO ID NOS:34, 35 and 36. Based on the biological assays described below, the preferred mature human artemin consists of SEQ ID NO:3 and similarly, the preferred mature mouse artemin consists of SEQ ID NO:34, respectively. The preferred human proregion polypeptide would thus comprise amino acids 40-107 of Fig. 2B (SEQ ID NO:50) and the corresponding preferred mouse pro-region polypeptide would comprise amino acids 40-111 of Fig. 2B (SEQ ID NO:51). Similarly the preferred human pre-pro- region polypeptide would thus comprise amino acids 1-107 of Fig. 2B (SEQ ID NO:52) and the corresponding preferred mouse pre-pro- region polypeptide would comprise amino acids 1-111 of Fig. 2B (SEQ ID NO:53). The mature, secreted artemin molecule is likely to form a disulfide linked homodimer by analogy to other members of the TGF-ß family.

The human and mouse preferred pre- region polynucleotides comprise nucleotides 1-117 of Fig. 1B (SEQ ID NO:54) and nucleotides 1-117 of Fig. 1C (SEQ ID NO:55), respectively; the human and mouse pro- region polynucleotides comprise nucleotides 118-321 of Fig. 1B (SEQ ID NO:56) and nucleotides 118-333 of Fig. 1C (SEQ ID NO:57), respectively; and the human and mouse pre-pro-region polynucleotides comprise nucleotides 1-321 of Fig. 1B (SEQ ID NO:58) and nucleotides 1-333 of Fig. 1C (SEQ ID NO:59), respectively. It is noted, however, that

alternative start points and alternative non-consensus cleavage points as discussed above would result in coding sequences corresponding to the alternative isoforms of pre-pro-artemin and mature artemin.

A preferred artemin according to the present invention is prepared by recombinant DNA technology although it is believed that artemin can be isolated in purified form from cell-conditioned medium as was done for neurturin.

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By "pure form" or "purified form" or "substantially purified form" it is meant that an artemin composition is substantially free of other proteins which are not artemin. Preferably, a substantially purified artemin composition comprises at least about 50 percent artemin on a molar basis compared to total proteins or other macromolecular species present. More preferably, a substantially purified artemin composition will comprise at least about 80 to about 90 mole percent of the total protein or other macromolecular species present and still more preferably, at least about 95 mole percent or greater.

Recombinant artemin may be made by expressing the DNA sequences encoding artemin in a suitable transformed host cell. Using methods well known in the art, the DNA encoding artemin may be linked to an expression vector, transformed into a host cell and conditions established that are suitable for expression of artemin by the transformed cell.

Any suitable expression vector may be employed to produce recombinant artemin such as, for example, the mammalian expression vector pCB6 (Brewer, *Meth Cell Biol 43*:233-245, 1994) or the *E. coli* pET expression vectors, specifically, pET-30a (Studier et al., *Methods Enzymol 185*:60-89, 1990). Other suitable expression vectors for expression in mammalian and bacterial cells are known in the art as are expression vectors for use in yeast or insect cells. Baculovirus expression systems can also be employed.

A number of cell types may be suitable as host cells for expression of recombinant artemin. Mammalian host cells include, but are not limited to, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo 205 cells, 3T3 cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from *in vitro* culture of primary tissue, primary explants, HeLa cells, mouse L cells, BHK, HL-60, U937,

HaK and Jurkat cells. Yeast strains that may act as suitable host cells include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, and any other yeast strain capable of expressing heterologous proteins. Host bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium and any other bacterial strain capable of expressing heterologous proteins. If the polypeptide is made in yeast or bacteria, it may be necessary to modify the polypeptide, for example, by phosphorylation or glycosylation of the appropriate sites using known chemical or enzymatic methods, to obtain a biologically active polypeptide.

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The polypeptide of the invention can also be expressed in transgenic plants (see, for example, U.S. Patent No. 5,679,880) or transgenic animals such as, for example, cows, goats, pigs, or sheep whose somatic or germ cells contain a nucleotide sequence encoding human artemin.

The expressed artemin polypeptide can be purified using known purification procedures, such as gel filtration and ion exchange chromatography. Purification may also include affinity chromatography using an agent that will specifically bind the artemin polypeptide, such as a polyclonal or monoclonal antibody raised against a mature artemin or fragment thereof. Other affinity resins typically used in protein purification may also be used such as concanavalin A-agarose, heparin-toyopearl® or Cibacrom blue 3GA Sepharose®. Purification of artemin can also include one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether.

It is also contemplated that an artemin polypeptide may be expressed as a fusion protein to facilitate purification. Such fusion proteins, for example, include an artemin amino acid sequence fused to a histidine tag such as when expressed in the pET bacterial expression system as well as the artemin amino acid sequence fused to the amino acid sequence of maltose binding protein (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX). Similarly, the polypeptide of the invention can be tagged with a heterologous epitope and subsequently purified by immunoaffinity chromatography using an antibody that specifically binds such epitope. Kits for expression and purification of such fusion proteins and tagged proteins are commercially available.

Artemin and fragments thereof may also be produced by chemical synthesis using methods known to those skilled in the art.

Artemin may be expressed in the monomeric units or such monomeric form may be produced by preparation under reducing conditions. In such instances refolding and renaturation can be accomplished using one of the agents noted above that is known to promote dissociation/association of proteins. For example, the monomeric form can be incubated with dithiothreitol followed by incubation with oxidized glutathione disodium salt followed by incubation with a buffer containing a refolding agent such as urea.

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Because multiple RXXR cleavage sites are present in the pro-artemin sequence, it is believed that isoforms of mature artemin exist. In addition, alternative non-consensus cleavage sites might also result in different isoforms. Thus, the mature artemin polypeptide may have a variable number of amino acids preceding the first canonical cysteine. Such alternate cleavage sites could be utilized differently among different organisms and among different tissues of the same organism. The N-terminal amino acids preceding the first of the seven conserved cysteines in the mature forms of members of the TGF-ß family vary greatly in both length and sequence. Furthermore, insertion of a ten amino acid sequence two residues upstream of the first conserved cysteine does not affect the known biological activities of one family member, dorsalin (Basler et al., *Cell 73*:687-702, 1993). By analogy, it is believed that artemin proteins containing sequences of different lengths preceding the first canonical cysteine may exist or could be made and that these would retain their biological activity.

The present invention also encompasses isolated polynucleotides comprising nucleotide sequences that encode any of the human and mouse artemin polypeptides described herein. As used herein, a polynucleotide includes DNA and/or RNA and thus the nucleotide sequences recited in the Sequence Listing as DNA sequences also include the identical RNA sequences with uracil substituted for thymine residues. Nucleotide sequences included in the invention are those encoding the human and mouse pre-pro-artemin amino acid sequences shown in Figs. 1B and 1C, respectively, as well as nucleotide sequences encoding the variant human artemin protein shown in Fig. 1D and nucleotide sequences encoding human and mouse pro-artemin (SEQ ID

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NOS:40 and 41, respectively). Preferred polynucleotides which encode human prepro- and pro- artemin comprise SEQ ID NOS:24 and 42, respectively, and preferred polynucleotides encoding mouse pre-pro- and pro-artemin comprise SEQ ID NOS:27 and 43, respectively. Polynucleotides within the scope of this invention do not include isolated chromosomes.

The invention also includes polynucleotides comprising a nucleotide sequence that encodes a mature artemin polypeptide such as mature human and mouse polypeptides consisting of any of SEQ ID NOS:3-5and SEQ ID NOS:34-36, respectively. Such polynucleotides encoding mature artemin include the human nucleotide sequences set forth in SEQ ID NOS:6-8 and the mouse nucleotide sequences set forth in SEQ ID NOS:37-39. A particularly preferred polynucleotide encodes SEQ ID NO:3 and comprises SEQ ID NO:6.

It is understood by the skilled artisan that degenerate nucleotide sequences can encode the artemin amino acid sequences described herein and these are also intended to be included within the present invention. For example, SEQ ID NO:44 represents the artemin coding sequence found in one human cDNA clone isolated by the inventors herein which contains G rather than A at position 582, but which does not change the encoded amino acid sequence. Such degenerate nucleotide sequences include modifications of naturally-occurring sequences in which at least one codon is substituted with a corresponding redundant codon preferred by a given host cell, such as *E. coli* or insect cells, so as to improve expression of recombinant artemin therein.

The present invention also encompasses vectors comprising an expression regulatory element operably linked to any of the artemin-encoding nucleotide sequences included within the scope of the invention. This invention also includes host cells, of any variety, that have been transformed with such vectors.

In yet another embodiment, a polynucleotide which specifically hybridizes to a human artemin-encoding polynucleotide or to its complement is provided. Specific hybridization is defined herein as the formation of hybrids between a polynucleotide, including oligonucleotides, and a specific reference polynucleotide (e.g., a polynucleotide comprising a nucleotide sequence complementary to a nucleotide sequence encoding human artemin) wherein the polynucleotide preferentially hybridizes to the specific reference polynucleotide over other non-artemin

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polynucleotides. Specifically hybridizing oligonucleotides are typically at least 15 nucleotides in length and are preferably at least 17 to at least 20 nucleotides long. Other preferred lengths include at least 22 to at least 25 nucleotides. A polynucleotide that specifically hybridizes to a reference sequence can be of any length, for example, about 15 nucleotides up to about 100 nucleotides or up to about 1000 nucleotides or up to about 10,000 nucleotides or even greater. Specific hybridization is preferably done under high stringency conditions which, as well understood by those skilled in the art, can readily be determined by adjusting several factors during hybridization and during the washing procedure, including temperature, ionic strength, length of hybridization or washing times, and concentration of formamide (see for example, Sambrook, Fritsch and Maniatis., *Molecular Cloning: a Laboratory Manual, 2d Ed., Vols. 1-3,* Cold Spring Harbor Laboratory Press, Plainview N.Y. 11803, 1989)

The present invention also includes nucleic acid sequences which encode for artemin polypeptides that have survival or growth promoting activity and that preferentially bind anti-human or anti-mouse artemin antibodies over other antibodies that do not bind to human or mouse artemin.

Methods are also provided herein for producing artemin. Preparation can be by isolation from conditioned medium from a variety of cell types so long as the cell type produces artemin. A second and preferred method involves utilization of recombinant methods by isolating a nucleic acid sequence encoding artemin, cloning the sequence along with appropriate regulatory sequences into suitable vectors and cell types, and expressing the sequence to produce artemin.

On the basis of the structural similarities of artemin to the sequences of neurturin, GDNF and persephin, artemin would be expected to promote the survival and growth of neuronal as well as non-neuronal cells. As discussed above, GDNF, neurturin and persephin influence a broad spectrum of neuronal populations in the peripheral and central nervous systems. Moreover, all other growth factors isolated to date have been shown to act on many different cell types (for example see Scully and Otten, *Cell Biol Int 19*:459-469, 1005; Hefti, *Neurotrophic Factor Therapy 25*:1418-1435, 1994 which are incorporated by reference). As an example of the actions of neurotrophic factors on non-neuronal tissues, the prototypical neurotrophic factor.

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NGF, also acts upon mast cells to increase their number when injected into newborn rats (Aloe, *J Neuroimmunol 18*:1-12, 1988). In addition, mast cells express the trk receptor and respond to NGF such that NGF is a mast cell secretogogue and survival promoting factor (Horigome et al., *J Biol Chem 269*:2695-2707, 1994). Moreover, members of the TGF-ß superfamily act on many cell types of different function and embryologic origin.

Thus, it is likely that artemin will have trophic activity on a variety of different neuronal cells, both peripheral and central, as well as on non-neuronal cells. With respect to neuronal cells, artemin supports the survival of neurons from all peripheral ganglia examined thus far, including sympathetic neurons and neural crest and placodally-derived sensory neurons, and also supports the survival of at least one population of CNS neurons, i.e., dopaminergic midbrain neurons. In addition, the inventors herein have detected artemin expression in a number of adult and fetal tissues, including heart, kidney, lung, peripheral leukocytes and bone marrow, which further supports the conclusion that artemin can provide trophic support to a variety of neuronal and non-neuronal cells. This suggests a role for artemin in hematopoiesis, inflammation, allergy, and cardiomyopathy. Artemin's trophic activity on any particular target cell type can be determined by routine experimentation using standard reference models.

The invention also contemplates synthetic, pan-growth factors comprising at least one active domain of artemin combined with at least one active domain of one or more other non-artemin growth factors. (For example, see Ilag et al., *Proc Nat'l Acad Sci 92*:607-611, 1995). These pan-growth factors would be expected to have the combined activities or other advantageous properties of artemin and the one or more other growth factors. As such, these pan-growth factors are believed to be potent and multispecific growth factors that are useful in the treatment of a wide spectrum of degenerative diseases and conditions including conditions that can be treated by any or all of the parent factors from which the active domains were obtained. Such pangrowth factors might also provide synergistic effects beyond the activities of the parent factors (Barres et al., *supra*).

Pan-growth factors within the scope of the present invention can include chimeric or hybrid polypeptides that are constructed from portions of fragments of at

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WO 00/18799 PCT/US99/22604

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least two growth factors. Growth factors of the TGF-ß superfamily are structurally related having highly conserved sequence landmarks whereby family members are identified. In particular, seven canonical framework cysteine residues are nearly invariant in members of the superfamily (Kingsley, Genes & Dev 8:133-146, 1994) which is incorporated by reference). Chimeric polypeptide molecules can, therefore, be constructed from a sequence that is substantially identical to a portion of the artemin molecule, up to one or more crossover points, and one or more sequences each of which is substantially identical with a portion of another TGF-B superfamily member extending on the other side of the corresponding one or more crossover points. For example, a portion of the amino terminal end of the artemin polypeptide can be combined with a portion of the carboxy terminal end of a neurturin polypeptide or alternatively a portion of the amino terminal end of a neurturin polypeptide can be combined with a portion of the carboxy terminal end of an artemin polypeptide. Such portions of artemin or neurturin polypeptides are preferably from about 5 to about 95, more preferably from about 10 to about 90, still more preferably from about 20 to about 80 and most preferably from about 30 to about 70 contiguous amino acids and such portions of another, non-artemin or, as the case may be, non-neurturin TGF-B superfamily member are preferably from about 5 to about 95, more preferably from about 10 to about 90, still more preferably from about 20 to about 80 and most preferably from about 30 to about 70 contiguous amino acids. For example, a particular crossover point might be between the third and fourth canonical framework cysteine residues. The particular non-artemin TGF-ß family member could be selected from family members including but not limited to transforming growth factor-\(\beta\)1 (TGF\(\beta\)1), transforming growth factor-\(\beta\)2 (TGF\(\beta\)2), transforming growth factor-B3 (TGFB3), inhibin B A (INHBA), inhibin B B (INHBB), the nodal gene (NODAL), bone morphogenetic proteins 2 and 4 (BMP2 and BMP4), the Drosophila decapentaplegic gene (dpp), bone morphogenetic proteins 5-8 (BMP5, BMP6, BMP7 and BMP8), the *Drosophila* 60A gene family (60A), bone morphogenetic protein 3 (BMP3), the Vg1 gene, growth differentiation factors 1 and 3 (GDF1 and GDF3), dorsalin (drsln), inhibin  $\alpha$  (INH $\alpha$ ), the MIS gene (MIS), growth factor 9 (GDF-9), glial-derived neurotrophic growth factor (GDNF), neurturin (NTN) and persephin. Furthermore, additional crossover points can be used to incorporate any desired

number of artemin portions or fragments with portions or fragments of any one or more other family members.

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In constructing a particular chimeric molecule, the portions of artemin and portions of the other, non-artemin growth factor are amplified using PCR, mixed and used as template for a PCR reaction using the forward primer from one and the reverse primer from the other of the two component portions of the chimeric molecule. Thus, for example a forward and reverse primers are selected to amplify the portion of artemin from the beginning to the selected crossover point between the third and fourth canonical cysteine residues using a artemin-encoding plasmid as template. A forward primer with a 5' portion overlapping with the artemin sequence and a reverse primer are then used to amplify the portion of the other, non-artemin growth factor member of the TGF-ß superfamily from the corresponding crossover point through the 3' end using a plasmid template containing the coding sequence for the non-artemin TGF-ß family member. The products of the two PCR reactions are gel purified and mixed together and a PCR reaction performed. Using an aliquot of this reaction as template a PCR reaction is performed using the artemin forward primer and the reverse primer for the non-artemin growth factor. The product is then cloned into an expression vector for production of the chimeric molecule.

Chimeric growth factors would be expected to be effective in promoting the growth and development of cells and for use in preventing the atrophy, degeneration or death of cells, particular in neurons. The chimeric polypeptides may also act as receptor antagonists of one or both of the full length growth factors from which the chimeric polypeptide was constructed or as an antagonist of any other growth factor that acts at the same receptor or receptors.

The present invention also includes therapeutic or pharmaceutical compositions comprising an artemin polypeptide in an effective amount for providing trophic support to cells in patients with cellular degeneration or dysfunction and a method comprising administering a therapeutically effective amount of the artemin polypeptide to a cell *ex vivo* or *in vivo*. The term "trophic support" is used herein to mean that a growth factor such as artemin provides sufficient nourishment to a cell such that the cell maintains or recovers at least one or more of its normal functions.

The compositions and methods of the present invention are useful for treating a number of degenerative diseases and anaplastic diseases. Where the cellular degeneration, dysfunction or anaplasia involves neurons, the diseases include, but are not limited to peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease, Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, nervous system tumors such as neuroblastomas, multiple sclerosis, peripheral nerve trauma or injury, exposure to neurotoxins, metabolic diseases such as diabetes or renal dysfunctions and damage caused by infectious agents. In addition, artemin compositions can be used to treat enteric diseases such as idiopathic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers. If the cellular degeneration or dysfunction involves nonneuronal cells such as bone marrow cells, artemin may be useful in treating diseases including, but not limited to disorders of insufficient blood cells such as, for example, leukopenias including eosinopenia and/or basopenia, lymphopenia, monocytopenia, neutropenia, anemias, thrombocytopenia as well as an insufficiency of stem cells for any of the above. The cellular degeneration or dysfunction can also involve myocardial muscle cells in diseases such as cardiomyopathy and congestive heart failure. In addition small cell lung carcinoma can be treated with artemin polypeptide or polynucleotide compositions.

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Treatment of enteric diseases with artemin includes the treatment of enteric neuropathies. The enteric nervous system is a complex collection of nerves that control the function of the gastrointestinal system, including gastrointestinal motility. Initial clinical studies with the NT-3, have shown that this neurotrophic factor increases gastrointestinal motility in normal volunteers and in patients suffering from peripheral neuropathies. Similarly, it is believed that artemin as well as the other members of the GDNF/neurturin/persephin family of growth factors will also show activity on enteric neurons. As a result, it is believed that artemin will be useful in treating enteric neuropathies such as in patients suffering from severe idiopathic constipation as well as patients suffering from constipation associated with Parkinson's disease, spinal cord injury, use of opiate pain-killers, and the like.

Whether artemin would be effectivity in the treatment of a particular cell type or tissues can be readily determined by one skilled in the art using any of a variety of

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assays known in the art. For example, with respect to providing trophic support for cells, trophic factors can produce beneficial biochemical and morphological effects and, under some circumstances, a promote cell survival. With respect to neurons, it is known in the art that depriving a neuron of trophic support results in a decrease in metabolic activity, i.e., glucose uptake, RNA synthesis and protein synthesis, required for normal function and growth. Deckwerth and Johnson, J. Cell Biol. 123:1207-1222, 1993. Removal of trophic support also results in a reduction in size of the cell body of the neuron. Presumably as a consequence of the loss of the metabolic effects of trophic factors, trophic factor deprivation results in a decrease or cessation of process outgrowth and may result in retraction of neuronal processes. In addition to the requirement of trophic factor for these aspects of neuronal biology, the neuron may require the neurotrophic factor to maintain survival; thus, survival assays are a frequently used means to detect or quantitate the actions of a neurotrophic factor. However, trophic support can also be manifest as morphological, biochemical, and functional changes; independent of neuronal number or any effect on survival.

As discussed above, growth factors can produce a cell differentiation in addition to providing trophic support for cells. Thus, it is believed that artemin polypeptides and polynucleotides can be beneficially used to produce a differentiation of anaplastic cells such as cancer cells. In particular, artemin can be used to treat nervous system tumors such as neuroblastomas. In addition, small cell lung carcinomas are known to express RET. Hence, it is believed that artemin can also be used to treat small cell lung carcinomas.

It is also contemplated that the eliciting of trophic support and/or differentiation can be achieved by administering an artemin polypeptide along with a GFRα3 polypeptide or by administering an artemin polynucleotide and a GFRα3 polynucleotide using the methods described herein for administration of artemin polypeptides or artemin polynucleotides. As noted above, Mouse GFRα3 was first identified as an expressed sequence tag (EST) that is homologous to GFRa1 and GFRα2 (Baloh et al., 1998, supra; Jing et al., 1997, supra; Naveilhan et al., Proc Natl Acad Sci USA 95, 1295-300, 1998; Widenfalk et al., Eur. J. Neurosci. 10, 1508-1517, 1998; Worby et al., J Biol Chem 273, 3502-3508, 1998). Human GFRα3 was identified by using the human GFR \alpha 2 amino acid sequence as a query to search a

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database of Expressed Sequence Tags (dbEST database) with the BLAST search algorithm (Altschul et al. J.Mol.Biol. 215:403-410, 1990; see copending application entitled "GFRa3, A Novel Member of the GDNF Coreceptor Family" which is file concurrently with the present application on December 22, 1998 and which is incorporated in its entirety herein by reference). Of the EST sequences identified, several (AA049894, AA050083, AA041935, AA238748) did not correspond identically to either GFR $\alpha$ 1 or GFR $\alpha$ 2 but had significant homologies to both sequences. The clones corresponding to these EST's were acquired from the Washington University EST project and sequenced. One of these corresponded to a full-length mouse cDNA which the inventors herein named GFRa3. Sequence information from the mouse cDNA was used to identify human genomic and cDNA clones, and the corresponding human and mouse predicted protein sequences are shown in Figure 12. The predicted amino acid sequences for human and mouse GFRα3 indicate a ~38.8 kDa protein with a putative N-terminal signal sequence (amino acids 1-31 in hGFRα3 and 1-28 in mGFRα3 of Figure 12) (Nielsen et al., Protein Eng. 10:106, 1997), a mature GFRα3 polypeptide (amino acids 32-372 in hGFRα3 and 29-369 in mGFRα3 of Figure 12), three putative N-linked glycosylation sites, and a hydrophobic stretch of residues at the C-terminus consistent with a GPI signal peptide (amino acids 373-400 of hGFRα3 and 370-397 of mGFRα3 of Figure 12) (Undenfriend et al., Annu. Rev. Biochem. 64:563-591, 1995) like those present in the related proteins GFRα1 and GFRα2 (Klein et al., Nature 387: 717-721, 1997; Jing et al., Cell 85:1113-1124, 1996; Treanor et al., Nature 382:80-83, 1996; Baloh et al., Neuron 18:793-802, 1997.). The human nucleotide sequence encoding GFRα3 precursor (SEQ ID NO:65) is shown in Figure 13. This precursor sequence includes the encoding sequence for the N-terminal signal sequence (nucleotides 1-93 of Figure 13), the encoding sequence for mature GFR $\alpha$ 3 (nucleotides 94-1116 of Figure 13) and the encoding sequence for the C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide (nucleotides 1117-1203 of Figure 13).

As described below, artemin binds to  $GFR\alpha 3$  in the absence of RET and it is believed that the resulting ligand/coreceptor complex is capable of binding to and activating RET receptors expressed by a target cell. Thus, treatment with artemin and

WO 00/18799 PCT/US99/22604

37

GFR $\alpha$ 3 would be expected to increase the sensitivity of cells normally responsive to artemin and would also be expected to provide trophic support to cells that express RET but that are not normally responsive to artemin. Preferably, the artemin and GFR $\alpha$ 3 polypeptides are from the same species, i.e., human. It is also preferred that the GFR $\alpha$ 3 polypeptide be in soluble form, i.e., that it lack the GPI linkage to avoid potential undesirable interactions with cell membranes. As used herein a GFR $\alpha$ 3 polypeptide is intended to include the mature protein with or without the GPI anchor, as well as GFR $\alpha$ 3 fragments, particularly soluble fragments lacking a GPI anchor, that are capable of binding to both artemin and RET with such binding leading to the activation of RET.

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In certain circumstances, it may be desirable to modulate or decrease the amount of artemin expressed. Thus, in another aspect of the present invention, artemin anti-sense oligonucleotides can be made and a method utilized for diminishing the level of expression of artemin, respectively, by a cell comprising administering one or more artemin anti-sense oligonucleotides. By artemin anti-sense oligonucleotides reference is made to oligonucleotides that have a nucleotide sequence that interacts through base pairing with a specific complementary nucleic acid sequence involved in the expression of artemin such that the expression of artemin is reduced. Preferably, the specific nucleic acid sequence involved in the expression of artemin is a genomic DNA molecule or mRNA molecule that contains sequences of the artemin gene. Thus, the invention contemplates artemin anti-sense oligonucleotides that can base pair to flanking regions of the artemin gene, untranslated regions of artemin mRNA, the pre- or pro- portions of the artemin gene or the coding sequence for mature artemin protein. The term complementary to a nucleotide sequence in the context of artemin antisense oligonucleotides and methods therefor means sufficiently complementary to such a sequence as to allow hybridization to that sequence in a cell, i.e., under physiological conditions. The artemin antisense-oligonucleotides preferably comprise a sequence containing from about 8 to about 100 nucleotides and more preferably the artemin antisense oligonucleotides comprise from about 15 to about 30 nucleotides. The artemin antisense oligonucleotides can also contain a variety of modifications that confer resistance to nucleolytic degradation such as, for example, modified internucleoside

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linkages (Uhlmann and Peyman, *Chemical Reviews 90*:543-548, 1990; Schneider and Banner, *Tetrahedron Lett 31*:335, 1990), modified nucleic acid bases and/or sugars and the like.

The therapeutic or pharmaceutical compositions of the present invention can be administered by any suitable route known in the art including for example intravenous, subcutaneous, intramuscular, transdermal, intrathecal or intracerebral. Administration can be either rapid as by injection or over a period of time as by slow infusion or administration of slow release formulation. For treating tissues in the central nervous system, administration can be by injection or infusion into the cerebrospinal fluid (CSF). When it is intended that artemin be administered to cells in the central nervous system, administration can be with one or more agents capable of promoting penetration of artemin across the blood-brain barrier.

Artemin can also be linked or conjugated with agents that provide desirable pharmaceutical or pharmacodynamic properties. For example, artemin can be coupled to any substance known in the art to promote penetration or transport across the blood-brain barrier such as an antibody to the transferrin receptor, and administered by intravenous injection. (See for example, Friden et al., *Science 259*:373-377, 1993). Furthermore, artemin can be stably linked to a polymer such as polyethylene glycol to obtain desirable properties of solubility, stability, half-life and other pharmaceutically advantageous properties. (See for example Davis et al. *Enzyme Eng 4:*169-73, 1978; Burnham, *Am J Hosp Pharm 51*:210-218, 1994). Preferably, artemin is administered with a carrier such as liposomes or polymers containing a targeting moiety to limit delivery of artemin to targeted cells. Examples of targeting moieties include but are not limited to antibodies, ligands or receptors to specific cell surface molecules.

For nonparenteral administration, the compositions can also include absorption enhancers which increase the pore size of the mucosal membrane. Such absorption enhancers include sodium deoxycholate, sodium glycocholate, dimethyl-β-cyclodextrin, lauroyl-1-lysophosphatidylcholine and other substances having structural similarities to the phospholipid domains of the mucosal membrane.

The compositions are usually employed in the form of pharmaceutical preparations. Such preparations are made in a manner well known in the

WO 00/18799 PCT/US99/22604

pharmaceutical art. One preferred preparation utilizes a vehicle of physiological saline solution, but it is contemplated that other pharmaceutically acceptable carriers such as physiological concentrations of other non-toxic salts, five percent aqueous glucose solution, sterile water or the like may also be used. It may also be desirable that a suitable buffer be present in the composition. Such solutions can, if desired, be lyophilized and stored in a sterile ampoule ready for reconstitution by the addition of sterile water for ready injection. The primary solvent can be aqueous or alternatively non-aqueous. Artemin can also be incorporated into a solid or semi-solid biologically compatible matrix which can be implanted into tissues requiring treatment.

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The carrier can also contain other pharmaceutically-acceptable excipients for modifying or maintaining the pH, osmolarity, viscosity, clarity, color, sterility, stability, rate of dissolution, or odor of the formulation. Similarly, the carrier may contain still other pharmaceutically-acceptable excipients for modifying or maintaining release or absorption or penetration across the blood-brain barrier. Such excipients are those substances usually and customarily employed to formulate dosages for parenteral administration in either unit dosage or multi-dose form or for direct infusion into the cerebrospinal fluid by continuous or periodic infusion.

Dose administration can be repeated depending upon the pharmacokinetic parameters of the dosage formulation and the route of administration used.

It is also contemplated that certain formulations containing artemin are to be administered orally. Such formulations are preferably encapsulated and formulated with suitable carriers in solid dosage forms. Some examples of suitable carriers, excipients, and diluents include lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, calcium silicate, microcrystalline cellulose, polyvinylpyrrolidone, cellulose, gelatin, syrup, methyl cellulose, methyl- and propylhydroxybenzoates, talc, magnesium, stearate, water, mineral oil, and the like. The formulations can additionally include lubricating agents, wetting agents, emulsifying and suspending agents, preserving agents, sweetening agents or flavoring agents. The compositions may be formulated so as to provide rapid, sustained, or delayed release of the active ingredients after administration to the patient by employing procedures well known in the art. The formulations can also contain

substances that diminish proteolytic degradation and promote absorption such as, for example, surface active agents.

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The specific dose is calculated according to the approximate body weight or body surface area of the patient or the volume of body space to be occupied. The dose will also be calculated dependent upon the particular route of administration selected. Further refinement of the calculations necessary to determine the appropriate dosage for treatment is routinely made by those of ordinary skill in the art. Such calculations can be made without undue experimentation by one skilled in the art based on the activity of artemin for a particular cell type *in vitro*. The activity of artemin on various peripheral and central neurons in culture is described below and artemin activity on a particular target cell type can be determined by routine experimentation. Exact dosages are determined in conjunction with standard doseresponse studies. It will be understood that the amount of the composition actually administered will be determined by a practitioner, in the light of the relevant circumstances including the condition or conditions to be treated, the choice of composition to be administered, the age, weight, and response of the individual patient, the severity of the patient's symptoms, and the chosen route of administration.

In one embodiment of this invention, artemin may be therapeutically administered by implanting into patients vectors or cells capable of producing a biologically-active form of artemin or a precursor of artemin, i.e. a molecule that can be readily converted to a biological-active form of artemin by the body. In one approach cells that secrete artemin may be encapsulated into semipermeable membranes for implantation into a patient. The cells can be cells that normally express artemin or a precursor of artemin or the cells can be transformed to express artemin or a precursor thereof. In some embodiments, the cells are transformed to express and secrete both artemin and GFR $\alpha$ 3, preferably in a soluble form. It is preferred that the cell be of human origin and that the artemin be human artemin when the patient is human. However, the formulations and methods herein can be used for veterinary as well as human applications and the term "patient" as used herein is intended to include human and veterinary patients.

Cells can be grown *ex vivo* for use in transplantation or engraftment into patients (Muench et al., *Leuk & Lymph 16*:1-11, 1994). In another embodiment of the

present invention, artemin can be used to promote the *ex vivo* expansion of a cells for transplantation or engraftment. Current methods have used bioreactor culture systems containing factors such as erythropoietin, colony stimulating factors, stem cell factor, and interleukins to expand hematopoietic progenitor cells for erythrocytes, monocytes, neutrophils, and lymphocytes (Verfaillie, *Stem Cells 12*:466-476, 1994). These stem cells can be isolated from the marrow of human donors, from human peripheral blood, or from umbilical cord blood cells. The expanded blood cells are used to treat patients who lack these cells as a result of specific disease conditions or as a result of high dose chemotherapy for treatment of malignancy (George, *Stem Cells 12(Suppl 1):249-255*, 1994). In the case of cell transplant after chemotherapy, autologous transplants can be performed by removing bone marrow cells before chemotherapy, expanding the cells *ex vivo* using methods that also function to purge malignant cells, and transplanting the expanded cells back into the patient following chemotherapy (for review see Rummel and Van Zant, *J Hematotherapy 3:*213-218, 1994).

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It is also believed that artemin with or without GFR $\alpha$ 3 can be used for the *ex vivo* expansion of precursor cells in the nervous system. Transplant or engraftment of cells is currently being explored as a therapy for diseases in which certain populations of neurons are lost due to degeneration such as, for example, in Parkinson's disease (Bjorklund, *Curr Opin Neurobiol 2*:683-689, 1992). Neuronal precursor cells can be obtained from animal or human donors or from human fetal tissue and then expanded in culture using artemin. These cells can then be engrafted into patients where they would function to replace some of the cells lost due to degeneration. Because neurotrophins have been shown to be capable of stimulating the survival and proliferation of neuronal precursor cells such as, for example, NT-3 stimulation of sympathetic neuroblast cells (Birren et al., *Develop 119*:597-610, 1993), artemin could also function in similar ways during the development of the nervous system and could be useful in the *ex vivo* expansion of neuronal cells.

In a number of circumstances it would be desirable to determine the levels of artemin or soluble GFR $\alpha$ 3 in a patient. A change in the amount of endogenously produced artemin or GFR $\alpha$ 3 may play a role in certain disease conditions, particularly where there is cellular degeneration such as in neurodegenerative conditions or

diseases. Other neurotrophic factors are known to change during disease conditions. For example, in multiple sclerosis, levels of NGF protein in the cerebrospinal fluid are increased during acute phases of the disease (Bracci-Laudiero et al., *Neuroscience Lett 147*:9-12, 1992) and in systemic lupus erythematosus there is a correlation

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between inflammatory episodes and NGF levels in sera (Bracci-Laudiero et al.

NeuroReport 4:563-565, 1993).

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Thus, quantification of artemin levels may provide clinically useful information. Furthermore, in the treatment of degenerative conditions, compositions containing artemin can be administered and it would likely be desirable to achieve certain target levels of artemin in sera, in cerebrospinal fluid or in any desired tissue compartment. It would, therefore, be advantageous to be able to monitor the levels of artemin in a patient. Accordingly, the present invention also provides methods for detecting the presence of artemin in a sample from a patient.

The term "detection" as used herein in the context of detecting the presence of artemin in a patient is intended to include the determining of the amount of artemin or the ability to express an amount of artemin in a patient, the distinguishing of artemin from other growth factors, the estimation of prognosis in terms of probable outcome of a degenerative disease and prospect for recovery, the monitoring of the artemin levels over a period of time as a measure of status of the condition, and the monitoring of artemin levels for determining a preferred therapeutic regimen for the patient.

To detect the presence of artemin in a patient, a sample is obtained from the patient. The sample can be a tissue biopsy sample or a sample of blood, plasma, serum, CSF or the like. Samples for detecting artemin can be taken from any tissue known to express artemin. When assessing peripheral levels of artemin, it is preferred that the sample be a sample of blood, plasma or serum or alternatively from a tissue biopsy sample. When assessing the levels of artemin in the central nervous system a preferred sample is a sample obtained from cerebrospinal fluid.

In some instances it is desirable to determine whether the artemin gene is intact in the patient or in a tissue or cell line within the patient. By an intact artemin gene it is meant that there are no alterations in the gene such as point mutations, deletions, insertions, chromosomal breakage, chromosomal rearrangements and the

WO 00/18799 PCT/US99/22604

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like wherein such alteration might alter production of artemin or alter its biological activity, stability or the like to lead to disease processes or susceptibility to cellular degenerative conditions. Conversely, by a non-intact artemin gene it is meant that such alterations are present. Thus, in one embodiment of the present invention a method is provided for detecting and characterizing any alterations in the artemin gene. The method comprises providing a polynucleotide that specifically hybridizes to an artemin cDNA, genomic DNA or a fragment thereof.

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Typically, patient genomic DNA is isolated from a cell sample from the patient and digested with one or more restriction endonucleases such as, for example, TaqI and AluI. Using the Southern blot protocol, which is well known in the art, this assay determines whether a patient or a particular tissue in a patient has an intact artemin gene or an abnormality in the artemin gene. Hybridization to the artemin gene would involve denaturing the chromosomal DNA to obtain a single-stranded DNA; contacting the single-stranded DNA with a gene probe associated with the artemin gene sequence; and identifying the hybridized DNA-probe to detect chromosomal DNA containing at least a portion of the human artemin gene.

The term "probe" as used herein refers to a structure comprised of a polynucleotide which forms a hybrid structure with a target sequence, due to complementarity of probe sequence with a sequence in the target region. The probes need not contain the exact complement of the target sequence, but must be sufficiently complementary to selectively hybridize with the strand being detected. By selective hybridization or specific hybridization it is meant that a polynucleotide preferentially hybridizes to a target polynucleotide. Oligomers suitable for use as probes may contain a minimum of about 8-12 contiguous nucleotides which are complementary to the targeted sequence and preferably a minimum of about 15 or 17 nucleotides although polynucleotide probes of about 20 to 25 nucleotides and up to about 100 nucleotides or even greater are within the scope of this invention.

The artemin gene probes of the present invention can be DNA or RNA oligonucleotides and can be made by any method known in the art such as, for example, excision, transcription or chemical synthesis. Probes may be labeled with any detectable label known in the art such as, for example, radioactive or fluorescent labels or enzymatic marker. Labeling of the probe can be accomplished by any

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method known in the art such as by PCR, random priming, end labeling, nick translation or the like. One skilled in the art will also recognize that other methods not employing a labeled probe can be used to determine the hybridization. Examples of methods that can be used for detecting hybridization include Southern blotting, fluorescence *in situ* hybridization, and single-strand conformation polymorphism with PCR amplification.

Hybridization is typically carried out at 25-45°C, more preferably at 32-40°C and more preferably at 37-38°C. The time required for hybridization is from about 0.25 to about 96 hours, more preferably from about one to about 72 hours, and most preferably from about 4 to about 24 hours.

Artemin gene abnormalities can also be detected by using the PCR method or any other known DNA amplification method, which uses oligonucleotides to identify a target sequence within a longer sequence, and primers that flank or lie within the artemin gene. The PCR method is well known in the art. Briefly, this method is performed using two oligonucleotide primers which are capable of hybridizing to the nucleic acid sequences flanking a target sequence that lies within an artemin gene and amplifying the target sequence. The terms "oligonucleotide primer" as used herein refers to a short strand of DNA or RNA typically ranging in length from about 8 to about 30 bases. The upstream and downstream primers are preferably a minimum of from about 15 nucleotides to about 20 nucleotides and up to about 30 nucleotides or even greater in length. The primers can hybridize to the flanking regions for replication of the target nucleotide sequence. The polymerization is catalyzed by a DNA-polymerase in the presence of deoxynucleotide triphosphates or nucleotide analogs to produce double-stranded DNA molecules. The double strands are then separated by any denaturing method including physical, chemical or enzymatic. Commonly, the method of physical denaturation is used involving heating the nucleic acid, typically to temperatures from about 80°C to 105°C for times ranging from about 1 to about 10 minutes. The process is repeated for the desired number of cycles.

The primers are selected to be substantially complementary to the strand of DNA being amplified. Therefore, the primers need not reflect the exact sequence of the template, but must be sufficiently complementary to selectively hybridize or

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PCT/US99/22604

specifically hybridize with the strand being amplified. By selective hybridization or specific hybridization it is meant that a polynucleotide preferentially hybridizes to a target polynucleotide.

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After PCR amplification, the DNA sequence comprising an artemin-encoding nucleotide sequence or a fragment thereof is then directly sequenced and analyzed by comparison of the sequence with the sequences disclosed herein to identify alterations which might change activity or expression levels or the like.

In another embodiment a method for detecting artemin is provided based upon an analysis of tissue expressing the artemin gene. The method comprises hybridizing a polynucleotide probe to mRNA from a sample of tissues that normally express the artemin gene or from a cDNA produced from the mRNA of the sample. The sample is obtained from a patient suspected of having an abnormality in the artemin gene or from a particular patient tissue or cell type suspected of having an abnormality in the artemin gene.

To detect the presence of mRNA encoding artemin protein, a sample is obtained from a patient. The sample can be from blood or from a tissue biopsy sample. The sample may be treated to extract the mRNA contained therein. The resulting mRNA from the sample is subjected to gel electrophoresis or other size separation techniques.

The mRNA of the sample is contacted with a nucleic acid serving as a probe to form hybrid duplexes. The use of a labeled probes as discussed above allows detection of the resulting duplex.

High stringency conditions can be used in order to prevent false positives, that is hybridization to non-artemin nucleotide sequences. When using sequences that are not perfectly complementary to an artemin-encoding polynucleotide or a fragment thereof, less stringent conditions could be used, however, this would be a less preferred approach because of the likelihood of false positives. The stringency of hybridization is determined by a number of factors during hybridization and during the washing procedure, including temperature, ionic strength, length of time and concentration of formamide. These factors are outlined in, for example, Sambrook et al. (Sambrook, et al., 1989, *supra*).

In order to increase the sensitivity of the detection in a sample of mRNA encoding the artemin protein, the technique of reverse transcription/polymerization chain reaction (RT/PCR) can be used to amplify cDNA transcribed from mRNA encoding the artemin protein. The method of RT/PCR is well known in the art.

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Alternatively, an artemin target sequence in the reverse transcribed cDNA can be amplified and detected using any other known methodology such as ligase chain reaction methods, including gap LCR (G-LCR) and other variations, or self-sustained sequence replication (3SR) and its various modifications. In addition, the artemin mRNA can be detected directly by asymmetric gap LCR (AG-LCR). See, e.g., Leckie et al., "Infectious Disease Testing by Ligase Chain Reaction" in *Molecular Biology and Biotechnology*, R. A. Myers, ed., pp. 463-466, VCH Publishers, 1995.

The present invention further provides for methods to detect the presence of the artemin protein in a sample obtained from a patient. Any method known in the art for detecting proteins can be used. Such methods include, but are not limited to immunodiffusion, immunoelectrophoresis, immunochemical methods, binder-ligand assays, immunohistochemical techniques, agglutination and complement assays. (for example see *Basic and Clinical Immunology*, Sites and Terr, eds., Appleton & Lange, Norwalk, Conn. pp 217-262, 1991). Preferred are binder-ligand immunoassay methods including reacting antibodies with an epitope or epitopes of the artemin protein and competitively displacing the artemin protein.

Numerous competitive and non-competitive protein binding immunoassays are well known in the art. Antibodies employed in such assays may be unlabeled, for example as used in agglutination tests, or labeled for use in a wide variety of assay methods. Labels that can be used include radionuclides, enzymes, fluorescers, chemiluminescers, enzyme substrates or co-factors, enzyme inhibitors, particles, dyes and the like for use in radioimmunoassay (RIA), enzyme immunoassays, e.g., enzyme-linked immunosorbent assay (ELISA), fluorescent immunoassays and the like.

Polyclonal or monoclonal antibodies to the artemin protein or to an epitope

thereof can be made for use in immunoassays by any of a number of methods known
in the art. By epitope reference is made to an antigenic determinant of a polypeptide.

The term epitope can also include artemin-specific B cell epitopes or T helper cell

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WO 00/18799 PCT/US99/22604 47

epitopes. An epitope could comprise 3 amino acids in a spatial conformation which is unique to the epitope. Generally an epitope consists of at least 6 such amino acids. Methods of determining the spatial conformation of amino acids are known in the art, and include, for example, x-ray crystallography and 2 dimensional nuclear magnetic resonance.

One approach for preparing antibodies to a protein is the selection and preparation of an amino acid sequence of all or part of the protein, chemically synthesizing the sequence and injecting it into an appropriate animal, usually a rabbit or a mouse (See Example 10).

Oligopeptides can be selected as candidates for the production of an antibody to the artemin protein based upon the oligopeptides lying in hydrophilic regions, which are thus likely to be exposed in the mature protein.

Antibodies to artemin can also be raised against oligopeptides that include one or more of the conserved regions identified herein such that the antibody can crossreact with other family members. Such antibodies can be used to identify and isolate the other family members.

Methods for preparation of the artemin protein or an epitope thereof include. but are not limited to chemical synthesis, recombinant DNA techniques or isolation from biological samples. Chemical synthesis of a peptide can be performed, for example, by the classical Merrifeld method of solid phase peptide synthesis (Merrifeld, J Am Chem Soc 85:2149, 1963) or the FMOC strategy on a Rapid Automated Multiple Peptide Synthesis system (DuPont Company, Wilmington, DE) (Caprino and Han, *J Org Chem 37*:3404, 1972).

Polyclonal antibodies can be prepared by immunizing rabbits or other animals by injecting antigen followed by subsequent boosts at appropriate intervals. The animals are bled and sera assayed against purified artemin protein usually by ELISA or by bioassay based upon the ability to block the action of artemin. When using avian species, e.g. chicken, turkey and the like, the antibody can be isolated from the yolk of the egg. Monoclonal antibodies can be prepared after the method of Milstein and Kohler by fusing splenocytes from immunized mice with continuously replicating tumor cells such as myeloma or lymphoma cells. (Milstein and Kohler Nature 256:495-497, 1975; Gulfre and Milstein, Methods in Enzymology: Immunochemical

Techniques 73:1-46, Langone and Banatis eds., Academic Press, 1981). The hybridoma cells so formed are then cloned by limiting dilution methods and supernates assayed for antibody production by ELISA, RIA or bioassay.

WO 00/18799

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The unique ability of antibodies to recognize and specifically bind to target proteins provides an approach for treating an over expression of the protein. Thus, another aspect of the present invention provides for a method for preventing or treating diseases involving over expression of the artemin protein by treatment of a patient with specific antibodies to the artemin protein.

Specific antibodies, either polyclonal or monoclonal, to the artemin protein can be produced by any suitable method known in the art as discussed above. For example, murine or human monoclonal antibodies can be produced by hybridoma technology or, alternatively, the artemin protein, or an immunologically active fragment thereof, or an anti-idiotypic antibody, or fragment thereof can be administered to an animal to elicit the production of antibodies capable of recognizing and binding to the artemin protein. Such antibodies can be from any class of antibodies including, but not limited to IgG, IgA, IgM, IgD, and IgE or in the case of avian species, IgY and from any subclass of antibodies.

Preferred embodiments of the invention are described in the following examples. Other embodiments within the scope of the claims herein will be apparent to one skilled in the art from consideration of the specification or practice of the invention as disclosed herein. It is intended that the specification, together with the examples, be considered exemplary only, with the scope and spirit of the invention being indicated by the claims which follow the examples.

25 Example 1

This example illustrates the identification and cloning of polynucleotides encoding human and mouse artemin.

Artemin was discovered by using the full length murine pre-pro neurturin amino acid sequence (195 amino acids) to scan the High Throughout Genome Sequences (htgs) database. The database was searched using the tblastn feature of the BLAST 2.0.5 program using the default parameters (Altschul, et al., Nucleic Acids Res. 25:3389-3402, 1997). This search identified two human bacterial artificial

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chromosome (BAC) clones, AC005038 (NH0486I22) and AC005051 (RG037F03), with significant homology scores when aligned with murine pre-pro neurturin. When the BLAST search was repeated using the mature human neurturin sequence as a query, the same two BAC clones were identified. Using the sequences of these two BAC clones, two EST's (Expressed Sequence Tags) were also identified in the EST database using the blastn feature of BLAST 2.0.5 program using the default parameters. These ESTs were: AA931637 (oo35c12) from a lung carcinoma and AA533512 (nj96b11) from a prostate carcinoma. Analysis of these alignments indicated that portions of the anonymous htgs sequences possibly represented a new member of the GDNF-neurturin-persephin family. Indeed, although, the hgts sequences were later discovered to have numerous sequence errors, i.e. omissions, additions and incorrect bases, one of the BAC clones, AC005038, had a stretch of 197 nucleotides (nucleotides 67,464 to 67,660) which ultimately turned out to be identical to nucleotides 663-467 of the complementary strand of the artemin nucleic acid (Fig. 1B) and the other, AC005051, had a stretch of 183 nucleotides (nucleotides 113,379 to 113,561) identical to nucleotides 648-468 of the complementary strand of the artemin nucleic acid (Fig. 1B).

Because sequences in the hgts database were generally known to contain many sequence errors, oligonucleotide primers were designed from the sequences of the two htgs entries to obtain fragments of the gene for this new growth factor from human 20 genomic DNA. The primers were designed to amplify an approximate 708 nt fragment encoding a portion of the predicted new growth factor, which would extend from within the pro-domain, through the mature region of the protein, and end within the 3' untranslated region. Human genomic DNA was used as a template in a PCR reaction which utilized a forward primer M4206 [5'-25 TGGCCGCTCTGGCTGAGCA-3'] (SEQ ID NO:20) and reverse primer M4205 [5'-CGATCATCTAGACCACCGGTAAGGGTCCAGTCTGCAA-3'] (SEO ID NO:21). The PCR reaction was carried out using Klentag in Klentag buffer with additives (1.5 M betaine/1.5% DMSO; final concentration) and the following parameters: initial denaturation at 95°C for 5 min, then 95°C for 15 sec, 60°C for 30 30 sec, 68°C for 80 sec x 30 cycles. The amplified products were kinased with T4 polynucleotide kinase, the ends were blunted with E. coli DNA polymerase I (Klenow

fragment), and cloned into EcoRV digested BSKS plasmid. The nucleotide sequences of the resulting clones were obtained.

An additional, overlapping fragment was also amplified from human genomic DNA, which extends from the beginning of the mature region to the stop codon. This 5 was amplified by PCR using the forward primer M4207 (5'-TACAAGGACGATGACGATAAGGGGGCGCGGGGCTGCGCCTCGC AGCTG-3') (SEQ ID NO:22) and reverse primer M4205 (5'-CGATCATCTAGACCACCGGTAAGGGTCCAGTCTGCAA-3') (SEQ ID NO:23). The PCR reaction was carried out using Klentaq in Klentaq buffer with additives (5% DMSO; final concentration) and the following parameters: initial denaturation at 95°C 10 for 5 min, then 95°C for 15 sec, 60°C for 30 sec, 68°C for 80 sec x 30 cycles. The amplified products were kinased with T4 polynucleotide kinase, the ends were blunted with E. coli DNA polymerase I (Klenow fragment), and cloned into EcoRV digested BSKS plasmid. The nucleotide sequences of the resulting clones were 15 obtained.

The sequences from these fragments and the sequences of the entries in the htgs database were assembled using Seqman program (DNASTAR). The assembled nucleotide sequence, which is shown in Fig. 1A, encodes a partial pro-artemin amino acid sequence which extends from within the pro-domain, includes 3 potential RXXR cleavage sites and then includes a region corresponding to the mature artemin polypeptide, which is homologous to mature GDNF, neurturin and persephin as shown in Fig. 2A.

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Primers designed from this sequence were also used to PCR amplify a fragment of mouse genomic DNA which corresponded to the mouse artemin gene. This mouse DNA fragment was than used to screen mouse BAC genomic libraries and a BAC clone containing the mouse artemin gene was identified (data not shown).

Primers were made from the human and mouse artemin genomic sequences to perform rapid amplification of cDNA ends (RACE) PCR to obtain the 5' end of the artemin cDNA from several tissue libraries. PCR fragments were blunt-cloned into EcoRV digested BSKS as above, and sequenced entirely. Human RACE products were obtained from MARATHON RACE cDNA libraries (CLONTECH) prepared from pituitary, placenta, and kidney. Mouse RACE products were obtained from an

embryonic day 18 (E18) mouse MARATHON RACE library. Complete double-stranded sequence analysis of cloned PCR fragments from genomic DNA and cDNA was performed to generate contigs of the human and mouse cDNAs. At least two independent clones of each PCR fragment were sequenced. Genomic sequence of mouse BAC clone restriction fragments (subcloned into pBluescript), and sequences derived from portions of the human BAC clone, were used to confirm the cDNA sequences, which are shown in Fig. 1B (human) and Fig. 1C (mouse).

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Intron locations in the cDNA and splicing were confirmed by sequencing PCR amplified fragments from human and mouse cDNA libraries that cross both introns. As illustrated in Fig. 2C, the artemin gene has two introns. cDNAs containing a second starting methionine (Met\*) produced by alternative splicing of the first intron were also identified by RACE PCR from some human cDNA libraries; however, a similar message was not identified in mouse, and the mouse genomic sequence does not contain a methionine in this position. Mature artemin protein generated from either of the human messages would be identical.

Analysis of markers present on the artemin-encoding BAC clones indicated that the human artemin gene is located on chromosome 1p32-33, flanked by markers D1S190-artemin-D1S211 (telomeric to centromeric).

The human DNA sequence as shown in Figure 14, which includes the Artemin gene, was compiled from sequences from multiple independently cloned PCT products using human genomic DNA as template and from the human BAC clone sequences. The beginning of the sequence corresponds to nt 11472 of BAC clone AC005051, and nt 68862 of BAC clone AC005038 from the htgs database as described above.

A rat artemin cDNA fragment as shown in Figure 15 was generated using PCT from a rat Schwann cell cDNA library as template with the following primers: forward 5'-CCGGTGAGCGCTCTCGGCCT-3' (SEQ ID NO:76) and reverse primer 5'-TTCTGGATTCTCCCAGAGGAGTTC-3' (SEQ ID NO:77). PCT conditions were as follows: 95°C for 2 min; then 30 cycles of 95°C for 10 sec, 60°C for 30 sec; 72°C for 30 sec, then 72°C for 5 min;

WO 00/18799 PCT/US99/22604

# 52 Example 2

This example illustrates the expression of artemin in various human tissues.

As an initial survey of artemin expression in adult and embryonic human tissues, a human master RNA blot (MRB) (CLONTECH) containing normalized samples of poly(A)RNA was probed with a random hexamer <sup>32</sup>P-labeled fragment of the human artemin cDNA and signals were visualized using a PhosphorImager and (Molecular Dynamics). The results are shown in Fig. 5A.

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Relatively low level expression was observed in many adult tissues, with the highest level in the pituitary gland, placenta and trachea. Among human fetal tissues, kidney and lung showed the highest level of expression. While little signal was visible in the adult and fetal brain, low-level expression of artemin mRNA was observed in structures of the adult basal ganglia (subthalamic nucleus, putamen, substantia nigra) and in the adult thalamus, suggesting artemin may influence subcortical motor systems. Expression was also observed in the spinal cord, however the RNA preparation from this tissue contains DRG material as well as spinal cord, which likely contributes to the observed expression (see Figure 6).

To more precisely localize artemin expression during embryogenesis, a fragment of the rat artemin cDNA was generated using PCR and cloned into pBluescript. Sense and anti-sense <sup>33</sup>P-labeled RNA probes were generated from the cloned rat artemin cDNA fragment and used for *in situ* hybridization analysis of artemin mRNA expression in fresh-frozen embryonic day 14 (E14) rat embryos, which was performed as described previously (Araki et al., *Neuron 17*:353-361, 1998). The results are shown in Figs. 6A and 6B.

No signal was observed in the brain or spinal cord at this embryonic age. The most striking expression was observed in the nerve roots, but not the developing neurons, of the dorsal root ganglia (DRG) (Fig. 6A). Furthermore, significant diffuse expression was also observed surrounding the superior mesenteric artery, corresponding to either the surrounding mesenchyme or expression by cell of the artery itself (Fig. 6B). These data suggest the possibility that artemin may act as a survival/trophic factor for peripheral neurons, either in a paracrine fashion for developing sensory neurons of the DRG, or as a target-derived factor for autonomic innervation of the superior mesenteric artery. Interestingly, among potential GFRα

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receptors, expression of artemin is most complementary with that of the orphan GFRα3, which is expressed at high levels in peripheral ganglia, but which has no detectable expression in the developing CNS by *in situ* hybridization (Baloh et al., 1998, *supra*; Trupp et al., *Mol. Cell. Neurosci.*, 11:47-63, 1998; Widenfalk et al., *Eur. J. Neurosci.* 10:1508-1517, 1998; Worby et al., *J. Biol. Chem.* 271, 23619-23622, 1998). No signal was detected using the sense probe on any tissue samples.

The expression of artemin in the developing nerve roots at E14 suggested it is produced by Schwann cell precursors or immature Schwann cells. To investigate if Schwann cells express artemin, semi-quantitative RT-PCR analysis was performed on cDNA libraries prepared from primary cultures of Schwann cells isolated from early post-natal rats, from myelinating Schwann cells from the adult sciatic nerve and from Schwann cells isolated from the distal segment of the sciatic nerve at 16 hours, 3 days or 7 days following nerve transection. Cultures of purified Schwann cells from early post-natal rats was performed as described by Brockes et al., Brain Res. 165:105-118, 1979. Sciatic nerve transection in adult rats and generation of reverse-transcribed cDNA were performed as described elsewhere (Araki et al., supra; Baloh et al., Neuron 18:793-802, 1997). cDNA samples from the libraries were normalized to levels of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and then amplified by RT-PCR using as forward primer 5'-TCGCGACGGTGGCTCACCGGTCTT-3' (SEO ID NO:66) and as reverse primer 5'-GCACGAGCCGCTGCAGAAGCGGAA-3' (SEQ ID NO:67) and the following conditions: 95°C for 2 min followed by 36 cycles of 95°C, 20 sec; 54°C, 30 sec; and 72°C, 30 sec.; and then a final incubation at 72°C, 2 min. Products were separated on a 3% agarose gel and stained with Ethidium Bromide.

As shown in Fig. 6C, artemin is expressed at much higher levels in immature Schwann cells in culture than by mature myelinating Schwann cells (PS) of the adult sciatic nerve (N). However, artemin expression is up-regulated in the distal nerve segment after sciatic transection, a paradigm in which Schwann cells reacquire an immature state to support regenerating axons (reviewed in Scherer, *Curr. Opin Neurol.* 10:386-397, 1997). These data indicate that Schwann cells produce artemin, and furthermore suggest that artemin expression is regulated appropriately to influence developing and regenerating peripheral neurons.

#### Example 3

This example illustrates the production of recombinant artemin protein.

A PCR fragment was generated which corresponded to the mature human artemin coding sequence (Fig. 3A), with an NdeI site and an 8-histidine tag on the 5' end, and a KpnI site on the 3' end, and then cloned directly into the corresponding sites of the pET30a(+) bacterial expression vector (Novagen). The artemin-encoding expression vector was transformed into the *E. coli* strain BL21 and recombinant human artemin protein was produced and purified as described previously for neurturin (Creedon et al., *Proc. Natl. Acad. Sci. USA* 94:7018-7023, 1997).

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## Example 4

This example illustrates that artemin promotes the survival of peripheral neurons *in vitro*.

As shown above, expression of artemin suggests it may influence developing peripheral neurons, like GDNF and neurturin. To assess artemin's ability to support the survival of different peripheral neuronal populations *in vitro*, neurons from the dorsal root, trigeminal, nodose, and superior cervical ganglia (SCG) derived from postnatal day 1 (P1) Sprague-Dawley rats (Harlan Sprague-Dawley, IN) were cultured in the presence of artemin, GDNF, neurturin or persephin. Each GDNF ligand family member consisted of the human mature amino acid sequence and was produced by standard recombinant DNA methods. Recombinant GDNF, neurturin and persephin were obtained from Genentech while recombinant artemin was produced as described in Example 3.

The cultures were performed using standard methods described elsewhere (Kotzbauer et al., *Neuron* 12:763-773, 1994; Kotzbauer et al., *Nature* 384:467-470, 1996; Milbrandt et al., *Neuron* 20:245-253, 1998). In brief, for SCG neurons, after dissection and dissociation, neurons were plated on collagen-coated 24-well tissue culture plates and maintained in NGF-containing medium (AM50) for 5 days, after which they were either kept in AM50 or switched to medium containing neutralizing anti-NGF antibodies plus GDNF, neurturin, persephin, artemin, (50 ng/ml where not noted otherwise), or no growth factors. Cultures were maintained for 3 days, after

which they were fixed in 4% paraformaldehyde and stained with toluidine blue, and surviving neurons were counted. For nodose ganglion and dorsal root ganglion neuron cultures, dissociated neurons were plated directly into NGF (for dorsal root ganglia), or BDNF (for nodose ganglia, 100ng/ml) or one of the GDNF ligands in serum-containing medium (AM50 with anti-NGF antibodies), and cells surviving after 3 days in culture were counted as above. Trigeminal ganglia were dissected, dissociated, and plated directly into medium containing NGF or the indicated GDNF ligands, and the number of surviving cells was assessed after 3 days *in vitro*. For all culture systems, 2-3 independent experiments were performed. The results are shown in Figs. 7A-7E.

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Like GDNF and neurturin (NTN), artemin (ART) supported the survival of a subset of sensory neurons from both the dorsal root ganglion (DRG) and the trigeminal ganglion (TG), whereas persephin (PSP) did not (Figure 7A and 7C). Interestingly, in both these populations of sensory neurons ART supported a larger number of neurons than GDNF or neurturin, and supported a similar (DRG) or a greater number (TG) of neurons than NGF. Dose response analysis of artemin's survival promoting effect on DRG neurons (Fig. 7B) revealed an EC<sub>50</sub> of 1-3 ng/ml,  $\sim 10^{-10} M$ .

With respect to visceral sensory neurons of the nodose ganglion (NG), which consists of a large number of neurons responsive to the neurotrophin BDNF (Lindsay et al., 1985), artemin, GDNF, and neurturin supported similar numbers of NG neurons, and each factor yielded equal or greater survival promotion than BDNF (Fig. 7D). These results were similar to previous reports (Kotzbauer et al., 1996, *supra*; Trupp et al., 1995, *supra*).

Artemin also supported the survival of SCG neurons in culture (Fig. 7E), although fewer neurons were supported by artemin than by GDNF or neurturin. In contrast to the sensory neurons examined above, none of the GDNF family ligands supported as many neurons as NGF. Therefore, similar to GDNF and neurturin, and as consistent with its embryonic expression pattern, artemin is a survival factor for sensory and sympathetic peripheral neurons in culture.

WO 00/18799 PCT/US99/22604

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## Example 5

This example illustrates that RET-expressing neuroblastoma cell lines are responsive to artemin.

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Neuroblastoma cell lines are derivatives of peripheral sympathetic neuroblast tumors that respond to GDNF and neurturin and display a variety of GFRα/RET receptor profiles (Hishiki et al., *Cancer Res.* 58:2158-2165, 1998; Tansey et al., submitted). Artemin's ability to influence several of these neuronal cell lines was investigated to determine if responsive cells had a consistent receptor profile.

In one experiment, cells from the SH-SY5Y neuroblastoma cell line were plated at  $2x10^5$  cells/ml/well in 12-well tissue culture plates. After one day in culture cells were either left untreated (Cn) or stimulated with 50 ng/ml artemin (ART) or 10  $\mu$ M retinoic acid (RA), a known inducer of differentiation of SH-SY5Y cells. Three days after factor addition, differentiation was assessed and cells were photographed. As shown in Figs. 8A-8C, artemin and retinoic acid induced strong differentiation responses in the SH-SY5Y cell line as indicated by their robust neurite outgrowth and neuronal morphology.

In another experiment, the ability of artemin to induce proliferation of the NBL-S neuroblastoma cell line was examined. NBL-S cells were plated at  $5x10^4$  cells/well on 48-well plates in standard medium, either without treatment or in the presence of 50 ng/mL of GDNF, artemin, or persephin. Actively dividing cells undergoing DNA synthesis 30 hrs after factor addition were detected using the BrdU (colorimetric) cell proliferation assay kit according to the manufacturer's instructions (Boehringer-Mannheim) and the data are shown in Fig. 8D. Like GDNF, artemin stimulated proliferation of NBL-S cells, whereas persephin did not.

To confirm that RET was a required component of the artemin receptor, the responsiveness of the neuroblastoma cell lines CHP126, CHP134, and SAN, which do not express RET, were examined. These cell lines did not respond to GDNF, neurturin or artemin (data not shown). Because both the SH-SY5Y and NBL-S cell lines express GFR $\alpha$ 1, GFR $\alpha$ 2, and GFR $\alpha$ 3, as well as RET, these experiments did not suggest the usage of a particular coreceptor by artemin.

WO 00/18799 PCT/US99/22604

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#### Example 6

This example illustrates that artemin supports the survival of dopaminergic neurons.

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As shown above, artemin expression was not observed in the brain or spinal cord of embryonic rats at the age examined and was not observed in the fetal human brain, but was present at very low levels in some adult brain regions. Therefore artemin's ability to influence dopaminergic neurons from the rat embryonic ventral midbrain was compared to the survival promoting activity of the other GDNF ligand family members.

Embryonic day 14 ventral mesencephalon cultures were prepared by removing the entire mesencephalon into cold Leibovitz's L15 + 6 mg/ml glucose and dissecting the tissue while keeping it on ice. Following dissection, the tissue was digested in a mixture of dispase (1 mg/ml, Sigma) and collagenase (1 mg/ml, Worthington Biochemical) for 25 min. Tissue was then washed twice with modified N2 media and triturated 35 times. Cell density and viability was assessed using a hemocytometer to count trypan blue excluding cells. Cells were plated at 20,000 cells/well on 8-well chamber slides (coated with 125 ng/ml poly-D-lysine and 25 ng/ml laminin) in serum-free medium consisting of DME/Hams F12 (1:1), 1 mg/ml BSA, 5 μM insulin, 10 nM progesterone, 100 μM putrescine, 30 nM selenium, 10 ng/ml rat transferrin, 100 U/ml penicillin, 100 U/ml streptomycin. Factors were added within 15 min of plating. After 3 days in culture, the cells were fixed, stained for tyrosine hydroxylase, and the number of tyrosine hydroxylase staining (TH+) neurons were counted. The results from two independent experiments are shown in Fig. 7F as the percentage of surviving TH+ neurons over control.

Interestingly, artemin supported the survival of dopaminergic midbrain neurons, although there is no apparent artemin expression in the ventral midbrain at this age. Therefore, artemin can promote survival of central as well as peripheral neurons, and indicates that receptors for artemin are present in both these populations.

In summary, all of the biological responses of artemin in peripheral and central neurons, and in neuronal cell lines, suggest that artemin utilizes receptor components similar to or overlapping with GDNF and neurturin.

#### Example 7

This example illustrates that artemin signals through RET using the former orphan receptor  $GFR\alpha 3$  as a coreceptor.

Because artemin, GDNF and neurturin exhibit similar activity profiles for peripheral and central neuronal populations and because artemin's activity correlated with RET expression in neuroblastoma cell lines, artemin's ability to activate RET in primary cultured SCG neurons and in the artemin-responsive neuroblastoma cell line NBL-S was examined by performing Western blot analysis of Ret phosphorylation and MAP kinase activation as described previously (Baloh et al., 1997, *supra*; Creedon et al., 1997, *supra*).

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In brief, SCG neurons were dissected from postnatal day 1 (P1) rats and maintained in NGF-containing medium for 5 days, after which they were deprived of NGF by switching to NGF-free medium in the presence of anti-NGF antibodies.

After 2 hours without NGF, neurons were switched to medium containing NGF,

GDNF or artemin at 50 ng/ml for 20 minutes. Cells were washed in cold PBS, and collected in immunoprecipitation buffer (1 mM EDTA/1 mM EGTA/0.2mM NaVO<sub>3</sub>/1 mM Pefabloc/1 uM pepstatin A/10ug/ml leupeptin/2 ug/ml aprotinin/1% Triton X-100/0.5% Nonidet P-40/150 mM NaCl in 10 mM Tris, pH 7.4). The lysates were incubated with 30 μl agarose-conjugated anti-phosphotyrosine antibodies

(Calbiochem) at 4°C for 1 hr. Beads were then washed three times with

membranes. Membranes were blocked for 1 hr at 25°C in TBS containing 5% dry milk, incubated overnight 4°C with a 1:300 dilution of anti-Ret antibody (C-19; Santa Cruz), then washed (3X in TBS containing 0.1% Tween-20) and incubated in a 1:10,000 dilution of anti-rabbit antibodies conjugated to HRP (Jackson Immunoresearch). Membranes were washed 3X, incubated with SuperSignal ULTRA (Pierce) for 5 minutes, then exposed to film. For MAP kinase (MAPK) assays, a portion of the total lysate wars removed before immunoprecipitation, separated by

immunoprecipitation buffer, resuspended in SDS sample buffer, and boiled for 5 min.

Samples were separated using SDS-PAGE, and transferred to nitrocellulose

30 SDS-PAGE, and immunoblotted using an anti-phospho-MAP-kinase antibody (New England Biolabs).

NBL-S cells were plated at  $3x10^5$  cells/well in 6-well plates, and after 46 hr switched to low-serum (0.5%) medium for 2 hr, and then stimulated with NGF, GDNF or artemin (50 ng/ml) for 20 min. Cells were then collected and analyzed as above. For PI-PLC treatment, a parallel set of NBL-S cells were plated as above but treated with 500 mU/ml of PI-PLC (Boehringer-Mannheim) for 1 hr, and washed with PBS and stimulated with factors as above.

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The results are shown in Figs. 9A and 9B. Like GDNF, artemin induced tyrosine-phosphorylation of RET in SCG neurons and in the NBL-S cell line, and this activation was effective in eliciting downstream signaling as measured by activation of the MAP kinase pathway. Furthermore, pretreatment of the NBLS cell line with the enzyme PI-PLC, which specifically cleaves GPI-anchored proteins from the cell surface, abolished artemin's ability to activate RET and all downstream signaling, but as expected had no effect on NGF signaling (Figure 9B). Therefore, these data indicate that like the other members of the GDNF ligand family, artemin signals through the RET receptor tyrosine kinase, and requires a GPI-anchored coreceptor to do so.

To investigate which if any of the known coreceptor(s) artemin can utilize to activate RET, its ability to directly bind to soluble Fc-fusion forms of GFRα1, GFR $\alpha$ 2, or GFR $\alpha$ 3 was assessed. GFR $\alpha$ 1-Fc, GFR $\alpha$ 2-Fc and GFR $\alpha$ 3-Fc fusion proteins were obtained from R&D Systems. Binding assays were performed similarly to assays previously described (Sanicola et al., Proc. Natl. Acad. Sci., USA 94:6238-6243, 1997). Recombinant GDNF, neurturin, artemin, persephin, or purified bovine serum albumin (BSA; Pierce) were coated to Nunc-Immuno MaxiSorp microtiter plates at 325 ng/ml in TBS (10mM Tris-HCl pH 7.5, 150 mM NaCl) for 1 hour at 25°C. Plates were washed (3X in TBS/0.03% Tween-20), and then blocked (blocking solution: TBS/1% BSA) for 1 hr at 25°C. Receptor bodies diluted in blocking solution were added and incubated for 2 hr at 25°C, washed 5X, then incubated with 1:10,000 dilution of HRP-conjugated anti-human Fc antibodies in blocking solution (Jackson Immunoresearch) for 45 minutes at 25°C. Finally, wells were washed 5X and the presence of HRP was assayed by addition of the chromogenic substrate 3.3'. 5,5-tetramethylbenzidine for 5 min. The reaction was stopped by adding an equal volume of 0.5M H<sub>2</sub>SO<sub>4</sub>, and color was measured in a plate reader at 450 nm. Non-

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specific binding of receptor bodies to the plates was measured in BSA coated wells, and this binding was subtracted from all other measurements. All experiments were performed in duplicate and the results are shown in Figs. 9C-9E.

In this assay, GFR $\alpha$ 1-Fc was able to bind to both GDNF and neurturin with similar affinity, but not to persephin or artemin. GFR $\alpha$ 2-Fc bound to neurturin but not to GDNF or artemin, consistent with previous reports that GDNF can only bind to GFR $\alpha$ 2 in the presence of RET (Sanicola et al., 1997). Interestingly, the former orphan receptor GFR $\alpha$ 3-Fc was able to bind to artemin, but not to any of the other GDNF ligand family members, consistent with the inability of GFR $\alpha$ 3 and RET to form a functional receptor complex for GDNF, neurturin or persephin (Baloh et al., 1998, *supra*; Worby et al., 1998, *supra*). Each of the receptor bodies bound to their respective ligands with an apparent Kd of ~3nM, which is similar to the previously reported affinity between GFR $\alpha$ 1-Fc and GDNF in this assay (Sanicola et al., 1997). Therefore, artemin can bind to the former orphan receptor GFR $\alpha$ 3, with an affinity similar to GDNF and NTN binding to their respective receptors.

While the binding data above suggest that GFRα3/RET is the functional receptor for artemin, such direct binding studies alone have been unreliable in predicting all GDNF family receptor interactions because of the observation that RET can modulate receptor binding (Sanicola et al., 1997, *supra*). Thus, to directly test which receptor combinations can form functional receptors for artemin, expression plasmids for the individual GFRα coreceptors together with the Gal4-Elk/Gal4-Luc reporter system were transiently transformed into fibroblasts that stably express RET. This system, which utilizes the ability of the Gal4-Elk fusion protein to respond to MAP kinase activity and activate transcription of the Gal4-luciferase reporter, has been used previously to monitor NGF/TrkA activation of the MAP kinase pathway in PC12 cells (Vossler et al., *Cell* 89:73-82, 1997; York et al., *Nature* 392:622-626, 1998), and GDNF/RET activation of MAP kinase in neuroblastoma cell lines (Worby et al., 1998, *supra*; Worby et al., *J. Biol. Chem*.271:23619-23622, 1996).

Generation of the expression plasmids for rat GFR $\alpha$ 1 and human GFR $\alpha$ 2, as well as MG87 fibroblasts stably expressing human RET (RET-3T3), were described in detail previously (Baloh et al., 1997, *supra*; Creedon et al., 1997, *supra*). For generation of the GFR $\alpha$ 3 expression plasmid, a fragment containing the complete

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coding region of human GFR $\alpha$ 3 was PCR-amplified from a human pituitary cDNA library, and this fragment was directly cloned into the KpnI and BamHI sites of pCB6 (Brewer, *Methods in Cell Biol.* 43:233-245, 1994) using sites engineered into the PCR primers. The GAL4-Elk1 chimera expression plasmid was a gift of P. Stork (Vollum Institute, Oregon Health Sciences University).

NLF neuroblastoma cells, which naturally express GFRα2/RET (Tansey et al., submitted), or the RET-3T3 cells were plated at 85,000 cells/well in 12-well plates, and cotransfected with the reporter plasmids Gal4-Luc and CMV-Gal4-Elk (250 ng/well and 50 ng/well, respectively), CMV-lacZ (50 ng/well) for transfection normalization, one of the CMV-GFR\alpha expression plasmids (500 ng/well), and pBluescript (650 ng/well) as carrier for a total of 1.5µg DNA/well using the Superfect reagent (Qiagen) according to the manufacturer's instructions. RET-3T3 fibroblasts were exposed to DNA/Superfect mixtures at 37°C overnight, washed and placed in low-serum (0.5%) medium, and stimulated with 50 ng/ml GDNF, artemin or persephin for 6-8 hours before collection (48 hours post-transformation). NLF neuroblastoma cells were exposed to DNA/Superfect mixtures for 2 hours, placed in full-serum medium overnight for recovery, and then switched to low-serum (0.5%) medium containing 50 ng/m of GDNF, artemin or persephin for 24 hours before collection (48 hours post-transformation). Measurement of luciferase and β-galactosidase activity was measured using a luminometer and performed as described (Svaren et al., EMBO J 17:6010-6019, 1998). The average luciferase activity of duplicate samples was normalized to β-galactosidase activity of the cotransformed lacZ reporter, and fold activation was calculated by dividing the normalized activity in factor-treated cells by that in the no treatment control.

As seen in Fig. 10A, consistent with previous reports in multiple systems, GDNF activated RET signaling in cells expressing GFR $\alpha$ 1/RET or GFR $\alpha$ 2/RET, but not GFR $\alpha$ 3/RET (Baloh et al., 1998; Baloh et al., 1997; Jing et al., 1997; Sanicola et al., 1997; Suvanto et al., 1997; Worby et al., 1998). As predicted from the binding data, artemin was able to activate GFR $\alpha$ 3/RET, but not GFR $\alpha$ 2/RET receptor complexes. Interestingly, artemin was also able to activate RET in cells expressing GFR $\alpha$ 1/RET, although to a lesser degree than GDNF.

WO 00/18799 PCT/US99/22604

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These results were confirmed in the NLF cell line, which is a more neuronal cell line that natively responds to GDNF by RET and MAP kinase activation (unpublished data). As shown in Fig. 10B, NLF cells transformed with GFR $\alpha$ 1 responded more intensely to GDNF, and became responsive to artemin, confirming the results observed in fibroblasts. Furthermore, transformation of NLF cells with GFR $\alpha$ 3, but not GFR $\alpha$ 2, allowed the cells to respond to artemin stimulation. Transformation of NFL cells with GFR $\alpha$ 3 also decreased the ability of the cells to respond to GDNF, presumably because CMV-driven overexpression of GFR $\alpha$ 3 decreases the relative amount of GFR $\alpha$ 2/RET complexes on the cells, thereby decreasing the number of functional GDNF receptors.

In summary, direct binding data and *in vitro* receptor activation experiments together indicate that artemin is the only known GDNF family ligand for the  $GFR\alpha 3/RET$  receptor complex, but like GDNF and neurturin it can also activate the  $GFR\alpha 1/RET$  receptor complex.

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#### Example 8

This example illustrates the specificity and cross-talk in interactions of members of the GDNF ligand and  $GFR\alpha$  receptor families.

A schematic interaction diagram of the GDNF ligand family members and GFRα receptor family members is presented in Fig. 11. Similar to many other ligand/receptor systems, including the neurotrophins, the system is characterized by having preferred ligand/receptor pairs, however cross-talk between the different ligands and receptors is apparent. The addition of artemin to this diagram reveals several new features. First, both direct receptor binding experiments and receptor activation experiments presented herein indicate that artemin is the only GDNF family ligand capable of utilizing the GFRα3/RET receptor complex. One additional study indicated that GDNF was able bind to GFRα3 in the presence of RET (Trupp et al., 1998), however this interaction was of low-affinity in that it required cross-linking to be observed, and its relevance is unclear because it does not lead to RET activation in multiple experimental paradigms *in vitro* (Baloh et al., 1998; Trupp et al., 1998; Worby et al., 1998). Second, the GFRα1/RET receptor complex is highly

promiscuous, and although results from most in vitro paradigms agree that it is the preferred receptor for GDNF, it is believed that neurturin and artemin also utilize the GFRα1/RET receptor complex in some systems *in vivo*. Recent observations regarding differences in peripheral neuron losses between GDNF and GFRα1-deficient mice suggest that GDNF can use GFRα2/RET as a receptor *in vivo*, and confirm that alternative ligand/receptor interactions may have biological importance in the GDNF family (Cacalano et al., 1998; Enomoto et al., 1998). Furthermore, a recent paper analyzing neurotrophin knockout mice concluded that NT-3 can signal through TrkB *in vivo*, an interaction observed many years earlier *in vitro* but was thought to be irrelevant *in vivo* (Farinas I et al., 1998; Ip et al., 1993). Therefore, consideration of all possible ligand/receptor interactions identified *in vitro* is often necessary to understand results of *in vivo* analysis of neurotrophic factor influences.

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Results from direct binding and RET activation experiments presented herein further suggest that like the GDNF-GFR $\alpha2$  interaction, the artemin-GFR $\alpha1$  interaction appears dependent on the presence of RET, as direct binding of artemin to GFR $\alpha1$ -Fc, or GDNF to GFR $\alpha2$ -Fc receptor bodies was not observed (Fig. 9C; Sanicola et al., 1997). This may be due to the nature of the binding assay which utilizes soluble receptor bodies binding to immobilized ligand. However, this may alternatively reflect an additional level of specificity available to the GDNF ligand/GFR $\alpha$  system, in cases where the GFR $\alpha$  coreceptors are expressed in the absence of RET (Baloh et al., 1997; Golden et al., 1998; Trupp et al., 1997; Yu et al., 1998). In these situations (i.e. GFR $\alpha1$  in injured peripheral nerve, and GFR $\alpha2$  expression in cerebral cortex), where coreceptors are expressed in "trans" and are hypothesized to secrete or present ligand/coreceptor complexes to cells or axons expressing RET, only the RET-independent subset of binding interactions would be possible.

Deposit of Plasmid: The following plasmid has been deposited under the terms of the Budapest Treaty, with the American Type Culture Collection, 10801 University Blvd., Manassas, VA 20110-2209. The accession number indicated was assigned after successful verification of the presence of the plasmid in the deposit, and the requisite fees have been paid. Access to said plasmid will be available during pendency of the patent application to one determined by the Commissioner to be

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entitled thereto under 37 CFR 1.14 and 35 USC 122. All restriction on availability of said plasmid to the public will be irrevocably removed upon the granting of a patent based upon the application. Moreover, the designated deposits will be maintained for a period of thirty (30) years from the date of deposit, or for five (5) years after the last request for the deposit, or for the enforceable life of the U.S. patent, whichever is longer. Should the plasmid become nonviable or be inadvertently destroyed, it will be replaced with a viable plasmid. The deposited materials mentioned herein are intended for convenience only, and are not required to practice the present invention in view of the description herein, and in addition, these materials are incorporated herein by reference.

	Name of Plasmid	Deposit Date	ATCC No.
15	phART	December 22, 1998	

In view of the above, it will be seen that the several advantages of the invention are achieved and other advantageous results attained.

As various changes could be made in the above methods and compositions without departing from the scope of the invention, it is intended that all matter contained in the above description and shown in the accompanying drawings shall be interpreted as illustrative and not in a limiting sense.

All references cited in this specification, including patents and patent applications, are hereby incorporated by reference. The discussion of references herein is intended merely to summarize the assertions made by their authors and no admission is made that any reference constitutes prior art. Applicants reserve the right to challenge the accuracy and pertinency of the cited references.

### What is Claimed is:

- 1. An isolated and purified growth factor comprising an artemin amino acid sequence or a conservatively substituted variant thereof or a fragment thereof of at least 8 contiguous amino acids.
- 2. The isolated and purified growth factor of claim 1 which promotes survival of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosine-hydroxylase-expressing dopaminergic ventral midbrain neurons.
- 3. The isolated and purified growth factor of claim 1 comprising a mammalian sequence which is at least 75% identical to SEQ ID NO:19, SEQ ID NO:33 or a conservatively substituted variant thereof.
- 4. The isolated and purified growth factor of claim 3 comprising a human polypeptide sequence as set forth in SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO:5.
- 5. The isolated and purified growth factor of claim 4 comprising a human pro-artemin as set forth in SEQ ID NO:40 or a human pre-pro-artemin as set forth in SEQ ID NO:32 or a conservatively substituted variant thereof or a polypeptide comprising a non-artemin pre-pro- region and the human polypeptide sequence.
- 6. The isolated and purified growth factor of claim 3 comprising a mouse polypeptide sequence as set forth in SEQ ID NO:34, SEQ ID NO:35, SEQ ID NO:36 or a conservatively substituted variant thereof.
- 7. The isolated and purified growth factor of claim 6 comprising a mouse pro-artemin as set forth in SEQ ID NO:41 or a mouse pre-pro-artemin as set forth in SEQ ID NO:27 or a conservatively substituted variant thereof or a polypeptide comprising a non-artemin pre-pro- region and the mouse polypeptide.

- 8. The isolated and purified growth factor of claim 1 comprising an amino acid sequence identical to the artemin polypeptide encoded by the human cDNA contained in the DNA deposited with ATCC on December 22, 1998.
- 9. The isolated and purified growth factor of claim 1 comprising an artemin polypeptide produced by a process comprising the steps of:
- (a) transforming a host cell with human artemin cDNA clone deposited with ATCC on December 22, 1998 operably linked to expression regulatory elements and
  - (b) expressing artemin polypeptide encoded by the clone.
  - 10. An isolated and purified polypeptide comprising:
- (a) a pre- region of human artemin as set forth in SEQ ID NO:48 or a preregion of mouse artemin as set forth in SEQ ID NO:49,
- (b) a pro- region of human artemin as set forth in SEQ ID NO:50 or a proregion of mouse artemin as set forth in SEQ ID NO:51,
  - (c) a pre-pro- region of human artemin as set forth in SEQ ID NO:52 or a pre-pro- region of mouse artemin as set forth in SEQ ID NO:53, or
    - (d) a conservatively substituted variant of (a), (b) or (c)..
  - 11. A pan-growth factor comprising the artemin polypeptide fragment of claim 1 and a fragment of at least one other growth factor from the TGF- $\beta$  superfamily.
  - 12. A nucleic acid comprising a polynucleotide encoding the pan-growth factor of claim 11.
  - 13. A composition comprising the growth factor of claim 1 and a GFR $\alpha$  polypeptide.
  - 14. The composition of claim 13 wherein the GFR $\alpha$  polypeptide is a GFR $\alpha$ 3 polypeptide or a GFR $\alpha$ 1 polypeptide.

- 15. An isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto comprising a nucleotide sequence encoding a growth factor of claim 1 or a fragment of said nucleotide sequence consisting of at least 15 contiguous nucleotides.
- 16. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 15 comprising a nucleotide sequence encoding an artemin polypeptide which promotes survival of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosine-hydroxylase-expressing dopaminergic ventral midbrain neurons wherein said nucleic acid molecule specifically hybridizes to a mature human artemin nucleotide sequence as set forth in SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8 or to a mature mouse artemin nucleotide sequence as set forth in SEQ ID NO:37, SEQ ID NO:38 or SEQ ID NO:39.
  - 17. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 16 comprising a nucleotide sequence encoding an artemin polypeptide as set forth in SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:34, SEQ ID NO:35 or SEQ ID NO:36.
  - 18. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 17 comprising a nucleotide sequence as set forth in SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:37, SEQ ID NO:38, SEQ ID NO:39 or SEQ ID NO:44.
  - 19. A vector comprising expression regulatory elements operably linked to the nucleic acid molecule of claim 15.
    - 20. A host cell transformed with the vector of claim 19.
  - 21. The isolated and purified nucleic acid molecule of claim 15 comprising ATCC deposit made on December 22, 1998.

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- 22. A host cell transformed with the vector of claim 21.
- 23. The isolated and purified nucleic acid molecule of claim 15 comprising a polynucleotide encoding a polypeptide selected from the group consisting of a human pro-artemin as set forth in SEQ ID NO:41, a human pre-pro artemin as set forth in SEQ ID NO:26 or SEQ ID NO:32, a mouse pro-artemin as set forth in SEQ ID NO:42, a mouse pre-pro-artemin as set forth in SEQ ID NO:29 and a polypeptide comprising a non-artemin pre-pro- region sequence and a human or mouse mature artemin amino acid sequence.
- 24. The isolated and purified nucleic acid molecule of claim 23 comprising a human pro-artemin nucleotide as set forth in SEQ ID NO:42, a human pre-pro-artemin nucleotide as set forth in SEQ ID NO:24, SEQ ID NO:30 or SEQ ID NO:44, a mouse pro-artemin nucleotide as set forth in SEQ ID NO:43, or a mouse pre-pro-artemin nucleotide as set forth in SEQ ID NO:27.
- 25. A recombinant nucleic acid molecule comprising an artemin nucleotide sequence or complement thereto wherein the artemin nucleotide sequence encodes an amino acid sequence selected from the group consisting of a pre-pro-artemin polypeptide, a pro-artemin polypeptide, a mature artemin polypeptide, a conservatively substituted variant thereof and a fragment thereof having at least 8 contiguous amino acids.
- 26. The isolated and purified polynucleotide of claim 15 which is an artemin antisense oligonucleotide.
- 27. An isolated and purified nucleic acid molecule comprising a polynucleotide encoding:
- (a) a pre- region of artemin as set forth in SEQ ID NO:54 or SEQ ID NO:55;
- 5 (b) a pro- region of artemin as set forth in SEQ ID NO:56 or SEQ ID NO:57;

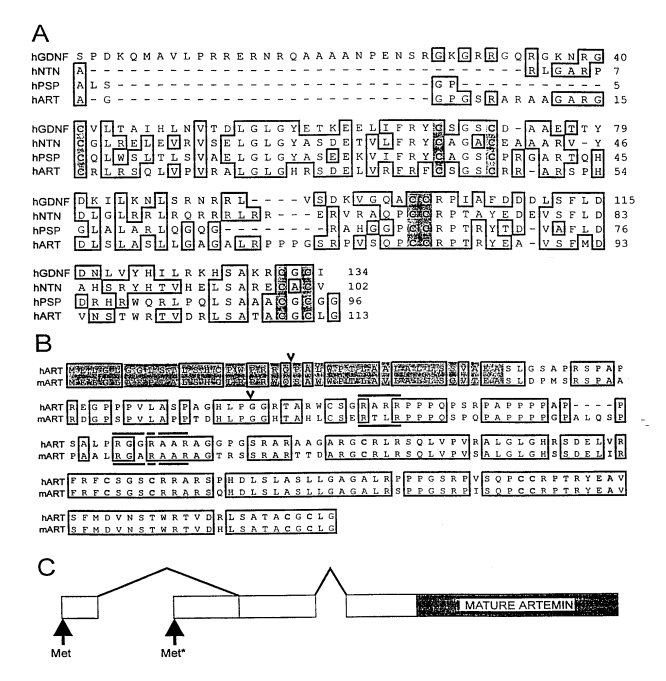
- (c) a pre-pro- region of artemin as set forth in SEQ ID NO:58 or SEQ ID NO:59; or
  - (d) a conservatively substituted variant of (a), (b) or (c).
- 28. An isolated and purified antibody which specifically reacts with the artemin polypeptide or fragment of claim 1.
- 29. A method for detecting expression of an artemin polypeptide in a sample comprising contacting the sample with an antibody according to claim 28 and detecting binding of the antibody to the artemin polypeptide.
- 30. A method for detecting expression of an artemin mRNA in a sample which comprises detecting a polynucleotide in the sample that specifically hybridizes to a polynucleotide consisting of SEQ ID NO:9.
- 31. The method of claim 30, wherein detecting the polynucleotide comprises:
- (a) contacting mRNA of the sample with a polynucleotide that specifically hybridizes to a polynucleotide consisting of SEQ ID NO:6; and
- (b) detecting the existence of a hybridization complex between the polynucleotide and the artemin mRNA.
  - 32. The method of claim 30, wherein the detecting step comprises:
- (a) producing a cDNA from the artemin mRNA using the reverse transcription method;
- (b) contacting the cDNA with at least two oligonucleotides that specifically hybridize to the cDNA to define a region of the cDNA to be amplified;
  - (c) amplifying the cDNA region; and
  - (d) detecting the amplified cDNA region.

- 33. A method for providing trophic support to and/or for producing differentiation of a cell comprising treating the cell with an effective amount of an artemin polypeptide or fragment thereof.
- 34. The method of claim 33, wherein the treating step comprises administering to the cell the artemin polypeptide or fragment with or without a  $GFR\alpha3$  polypeptide.
- 35. The method of claim 33, wherein the target cell is within a patient and the treating step comprises administering to the patient the artemin polypeptide or fragment with or without a GFR $\alpha$ 3 polypeptide.
- 36. The method of claim 33, wherein the target cell is within a patient and the treating step comprises administering to the patient a polynucleotide encoding the artemin polypeptide or fragment.
- 37. The method of claim 33, wherein the artemin polypeptide is expressed by a cell implanted into the patient.
- 38. The method of claim 33, wherein the target cell is a neuron in a patient suffering from peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease, Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, a nervous system tumor such as neuroblastomas, multiple sclerosis, infection or an enteric disease such as idiopathic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers or the target cell is a non-neuronal cell in a patient suffering from small cell lung carcinoma.

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GCGC	:CC	200	GG :	CGC R	GC(	CG(	CGC	CC	GA(	300	333 8	GGC	GCC P	CG G	TCC	GGC F	:GC R	ga A	GCC R	CG	TC	GCC A	G G	GGC A	CGC F	ccc {	CG₽ G	C rCd	GC R	GG2	ACG L	CG <i>i</i> R	AGC S	GT Ç	CG:	AC L	CA(	CGC	SCC P	ACC V	R R	:
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GCG7	AGO	CCG	<del>→  </del> GA	ccc	GG	TG	GCC	GAG	GC'	rgo	CTC	GA	CCA	ACG	CA	AA(	GC	GΑ	AGA	CC	TC	GCC	GAG	GG?	ACC	GC	GGG	CGC	:GC	GC	GAG	AG	<del>- -</del> GTC	TC	CT	GG	AG'	TC	GA	CCC	GGT	F 501
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TA	CGC	GG	CCG	G.	CT.	AG	AGI	CC	GG	CT	CC	TG	TC	GGC	GA	.GG	AA	CTC	CCA	GG	AAC	GA	GG	GG'	TTC	GG	GTO	GA	CCC	CAC	GG	SAG	AA.	AGA	AGG	GAC	TC	CGI	AGO	GTG/	AAC	CAG	
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C	AGC	GC	CT	CC	TG	CCC	SCC	GC	:GC	GC	GC'	rc'	rcc	CAC	AC	GA(	CCI	CP	GC	CT	GGC	CA	GC	CTA	ACT	GG	GCG	cc	GGG	GC	CCI	'GC	GAG	CCG	CC	ccc	GG	GCT	cc	CGG	GCC(	CGTC ···! GCAG	. 600
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FIGURE 2

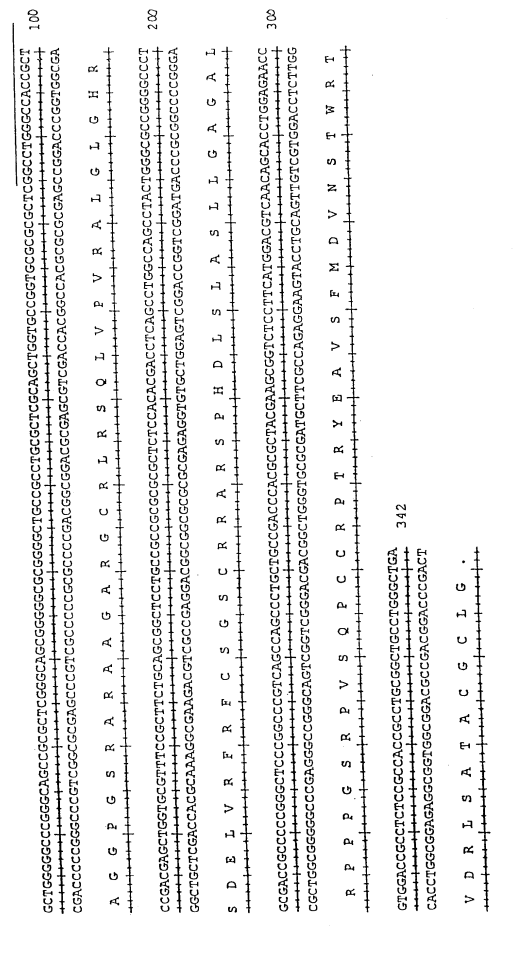


FIGURE 3A

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FIGURE 3B

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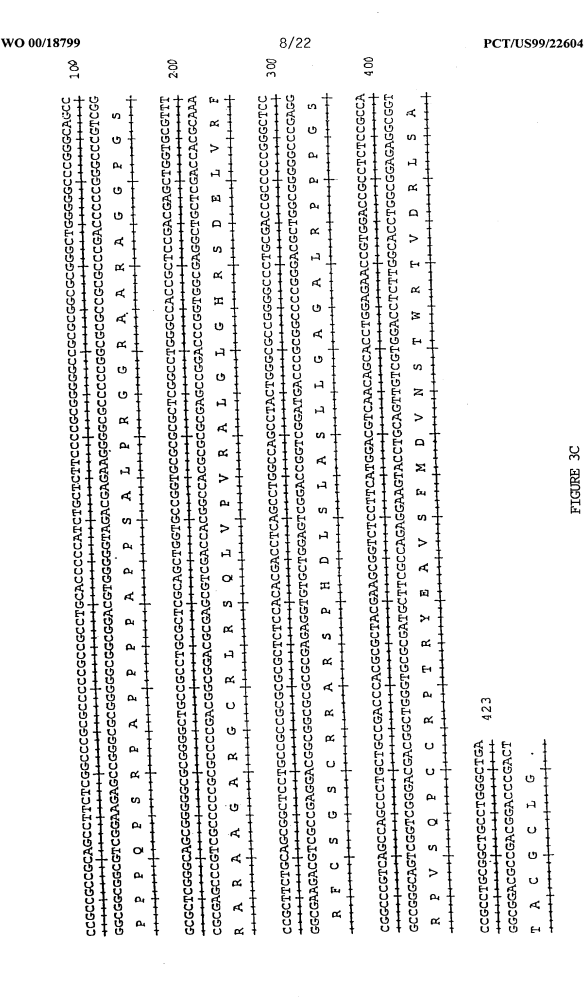
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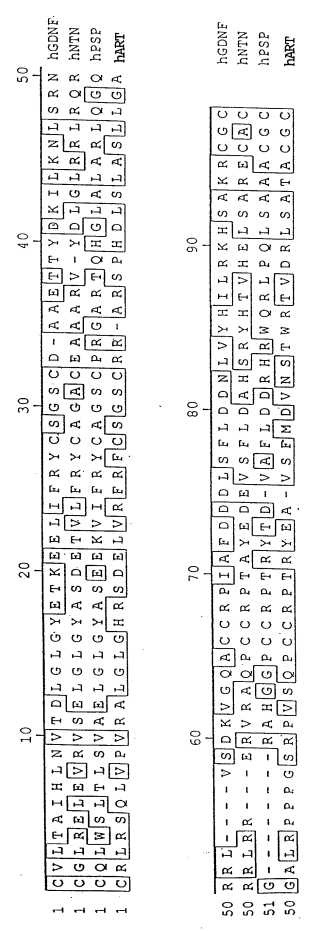
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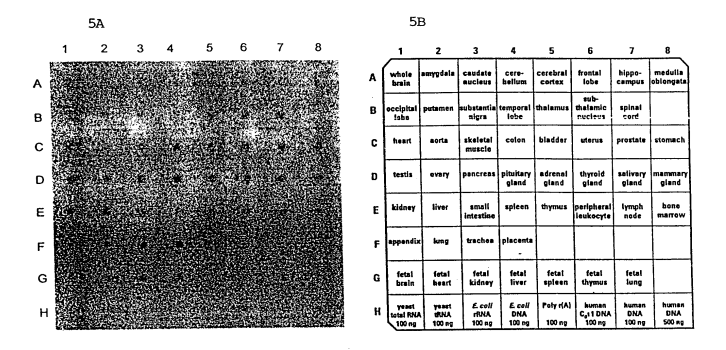


FIGURE 5

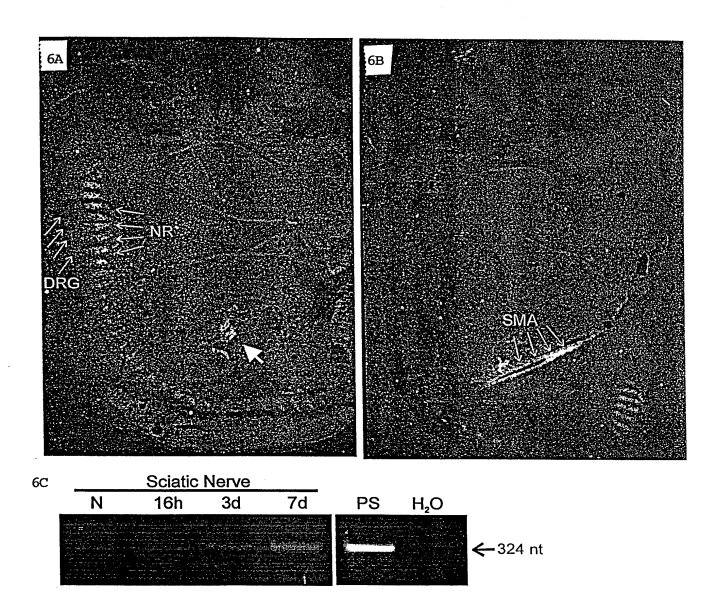


FIGURE 6

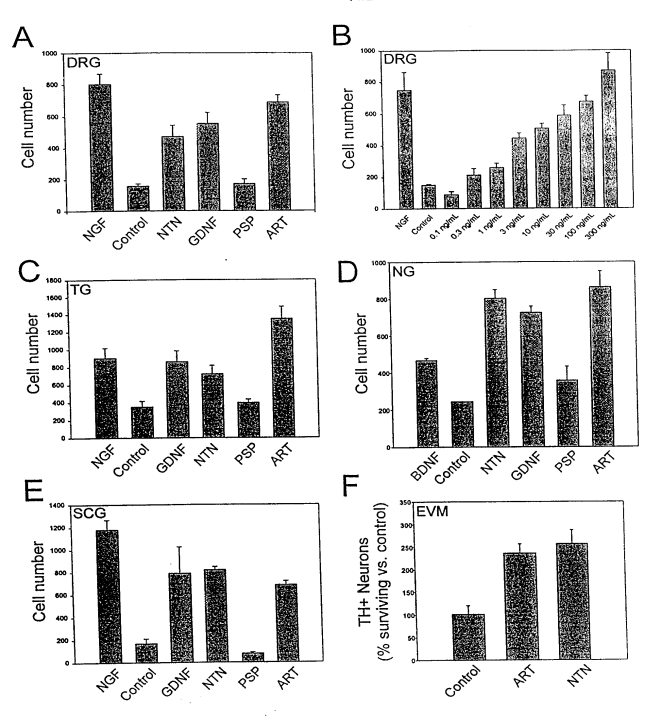


FIGURE 7

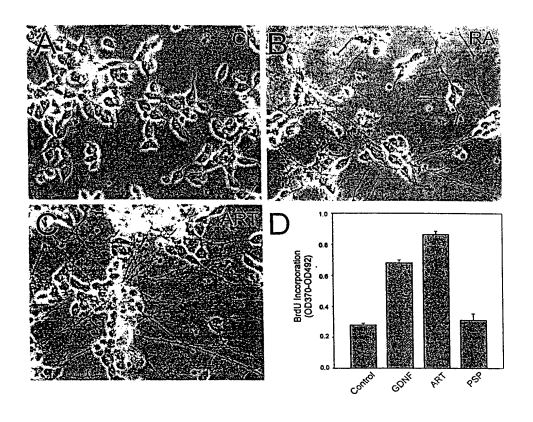


FIGURE 8

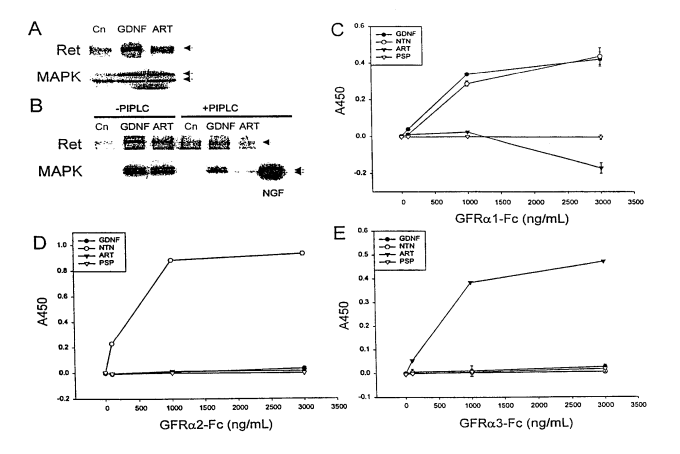


FIGURE 9

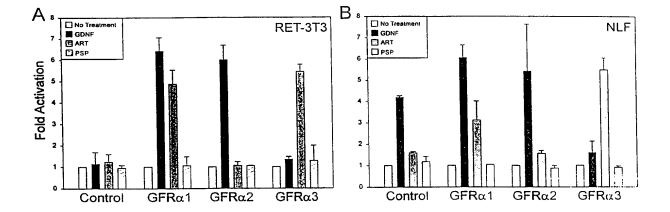


FIGURE 10

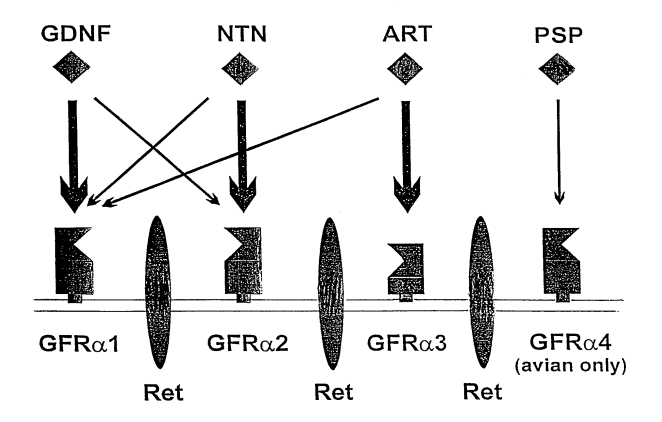


FIGURE 11

FIGURE 12

TTTCCAGACCCACTGCCATCCATGGACATCCTAGGAACTTGTGCAACAGAGCAGTCCAGATGTCTACGAGCATACCTGG AGTGGCAACCTGCAGGAGGGTGTGAAATGCTGGAAGGGTTCTTCTCCCACAACCCCTGCCTCACGGAGGCCATTGCAGC TCTCGCAGCCGGAGACCCCCTTCCCACAGAAAGCCGACTCATGAACAGCTGTCTCCAGGCCAGGAGGAAGTGCCAGGCTG TCGGTCCCTGCTGCTGCCTGGAGGCAGCACAGCAACTCAGGAACAGCTCTCTGATAGGCTGCATGTGCCACCGGCGCAT GAAGAACCAGGTTGCCTTGGACATCTATTGGACCGTTCACCGTGCCCGCAGCCTTGGTAACTATGAGCTGGATGTCT CCCCCTATGAAGACACAGTGACCAGCAAACCCTGGAAAATGAATCTCAGCAAACTGAACATGCTCAAACCAGACTCAGAC CTCTGCCTCAAGTTTGCCATGCTGTACTCTCAATGACAAGTGTGACCGGCTGCGCAAGGCCTACGGGGAGGCGTGCTC CTGCCGCCTGTGGCCCCCCAACTGCCTGCGGCGCCCTCTGCTTCTCCGACCCGCTTTGCAGATCACGCCTGGTGGA GGCTGATTGGGACTGCCATGACCCCCAACTTTGTCAGCAATGTCAACACCAGTGTTGCCTTAAGCTGCACCTGCCGAGGC TAAGATGCGTTTTCACAGCCAACTCTTCTCCCAGGACTGGCCACACCCTTTGCTTGTTGTTGGCACACCAGAATGAAA ACCCTGCTGTGAGGCCACAGCCCTGGGTGCCCTCTTTTCTCCTGCACGCTTCCCTTGATTCTGCTCCTGAGCCTATGG ATCCCACCTGCAGTGCTGCCTACCACCTGGATTCCTGCACCTCTAGCATAAGCACCCCCACTGCCTCAGAGGAGCCT

FIGURE 13

CTCTGAGCTTCTCTGAGCCTTGTTTGCTCATCTGGAAAAAGGGGATTAAACCATTTACCTCATGGAGTTGTGAAAGAATAGCTGCAAAGCACCTAACACA	100
GAGACTCGAAGAGACTCGGAACAACGAGTAGACCTTTTTCCCCTAATTTGGTAAATGGAGTACCTCAACACTTTCTTATCGACGTTTCGTGGATTGTGT  L . A S L S L V C S S G K R G L N H L P H G V V K E . L Q S T . H  S E L L . A L F A H L E K G D . T I Y L M E L . K N S C K A P N T  P L S F S E P C L L I W K K G I K P F T S W S C E R I A A K H L T H	
TAGTAAGGTTCCCAGTGCAGCTACTTCTGCTGGGTTGAGTCTAGCTGTGTAGGCCCCTTGTTCCTCACCTGGAGAAACTGGGGTGGCAGGCCGGTCCCCC ATCATTCCAAGGGTCACGTCGATGAAGACGACCCAACTCAGATCGACAACTCCGGGGAACAAGGAGTGGACCTCTTTGACCCCACCGTCCGGCCAGGGGG	200
I V R F P V Q L L L L G . V . L C R P L V P H L E K L G W Q A G P P G S Q C S Y F C W V E S S C V G P L F L T W R N W G G R P V P S K V P S A A T S A G L S L A V . A P C S S P G E T G V A G R S P	
ACAAAAGATAACTCATCTCTTAATTTGCAAGCTGCCTCAACAGGAGGGTGGGGGAACAGCTCAACAATGGCTGATGGGCGCTCCTGGTGTTGATAGAGAT	300
Q K I T H L L I C K L P Q Q E G G G T A Q Q W L M G A P G V D R D H K R . L I S . F A S C L N R R V G E Q L N N G . W A L L V L I E M T K D N S S L N L Q A A S T G G W G N S S T M A D G R S W C R	
GGAACTTGGACTTGGAGGCCTCTCCACGCTGTCCCACTGCCCTGGCCTAGGCGGCAGGTGAGTGGTTCTCCCAGTGACTCCTACCTGGTACTGAGGAAA	400
GTWTWRPLHAVPLPLA.AAGEWFSQ.LLPGTEE ELGLGGLSTLSHCPWPRRQVSGSPSDSYLVLRK WNLDLEASPRCPTAPGLGGR.VVLPVTPTWY.GK	
GGCGGCTTGACTGGTGAGGGAGGAGCAGGGCTTGGCTTG	500
R R L D W . G R A G L G L G S G . V W E G K W S G R D Q V N G R R S G G L T G E G E Q G L A W A A V R C G R E N G Q G G T R . M G G G A A . L V R E S R A W L G Q R L G V G G K M V R E G P G E W E E E	
CGGGACTTCTCTGAATGGTCGGTGCACTCAGGTGATTCCTCCCCTGGGCTCCCAGAGGCAGCAAACCCATTATACTGGAACCTAGGCCCTTCCTGAGTTT	600
GCCCTGAAGAGACTTACCAGCCACGTGAGTCCACTAAGGAGGGGACCCGAGGGTCTCCGTCGTTTGGGTAATATGACCTTGGATCCGGGAAGGACTCAAA	000
GTSLNGRCTQVIPPLGSQRQQTHYTGT.ALPEF AGLL.MVGALR.FLPWAPRGSKPIILEPRPFLSF RDFSEWSVHSGDSSPGLPEAANPLYWNLGPS.V	
CCCCTCCACACAGCTAGGAGCCCATGCCCGGCCTGATCTCAGCCCGAGGACAGCCCCTCCTTGAGGTCCTTCCT	700
PLHTARSPCPA.SQPEDSPSLRSFLPKPTWVPS PSTQLGAHARPDLSPRTAPP.GPSSPSPPGCPL SPPHS.EPMPGLISARGQPLLEVLPPQAHLGALF	
FIGURE 14A	

TCTCCCTGAGGCTCCACTTGGTCTCTCCGCGCAGCCTGGCCCACCCTGGCCGCTCTGGCTCTGAGCAGCGTCGCAGAGGCCTCCCTGGGC	200
AGAGGGACTCCGAGGTGAACCAGAGAGGCGCGTCGGACGGGACACCGGGTGGGACCGGGAGACCGAGACCGACGACTCCGCAGCGTCTCCGGAGGGACCCG	800
F S L R L H L V S P R S L P C G P P W P L W L C . A A S Q R P P W A S P . G S T W S L R A A C P V A H P G R S G S A E Q R R R G L P G L P E A P L G L S A Q P A L W P T L A A L A L L S S V A E A S L G	
TCCGCGCCCCGCAGCCCTGCCCCCGCGAAGGCCCCCCGCCTGTCCTGCCGTCCCCCGCCGGCCACCTGCCGGGTAGGTGAGAGGGGGGGG	900
PRPAALPPAKAPRLSWRPPPATCRVGERARGRG LRAPQPCPPRRPPACPGVPRRPPAG.VRGRGGGA SAPRSPAPREGPPPVLASPAGHLPGR.EGEGAG	
GGGGCTGGCCCGGGACACCGCGCGTGACTGGGTCTCATTCCAGGGGGACGCACGGCCCGCTGGTGCAGTGGAAGAGCCCGGCGGCGGCGCCGCCGCCGCCGCCGCCGCC	100
GAGPGHRA. LGLIPGGRTARWCSGRARRPPPQPGLARDTARDWVSFQGDARPAGAVEEPGGRRSLRGWPGTPRVTGSHSRGTHGPLVQWKSPAAAAAF	
CTCGGCCCGCCCCCCCCCCCCCCCCCCATCTGCTCTTCCCCGCGGGGGCCCGCGCGCG	110
SRPAPPPPAPPSALPRGGRAARAGGPGSRARAAG LGPRPRRLHPHLLFPAGAARRGLGARAALGQR SARAPAACTPICSSPRGPRGAGWGPGQPRSGSG	-
GGCGCGGGGCTGCCGCTCGCAGCTGGTGCCGGTGCGCGCGC	120
ARGCRLRSQLVPVRALGLGHRSDELVRFRFCSGGRGAAACARSWCRCARSAWATAPTSWCVSASAAAGAGLPPALAAGARARPGPPLRRAGAFPLLQR	
TCCTGCCGCCGCGCGCGCTCTCCACACGACCTCAGCCTGGCCAGCCTACTGGGCGCGGGGCCCTGCGACCGCCCCCGGGCTCCCGGCCCAGCCAG	130
S C R R A R S P H D L S L A S L L G A G A L R P P P G S R P V S Q P A A A R A L H T T S A W P A Y W A P G P C D R P R A P G P S A S L L P P R A L S T R P Q P G Q P T G R R G P A T A P G L P A R Q P A	
CCTGCTGCCGACCCACGCGCTACGAAGCGGTCTCCTTCATGGACGTCAACAGCACCTGGAGAACCGTGGACCGCCTCTCCGCCACCGCCTGCGGCTGCCT GGACGACGGCTGGGTGCGCGATGCTTCGCCAGAGGAAGTACCTGCAGTTGTCGTGGACCTCTTGGCACCTGGCGGAGAGGCGGTGGCGGACGACGGACG	. 140
P C C R P T R Y E A V S F M D V N S T W R T V D R L S A T A C G C L P A A D P R A T K R S P S W T S T A P G E P W T A S P P P P A A A L L P T H A L R S G L L H G R Q Q H L E N R G P P L R H R L R L P	
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TCCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGGCCACCACTACCTATAGTAGGGGCTTGTCCACTTCCCTGTTGACTGATCGTCGGGGTCTCGG  P G T K A S K L R G P C R W V M D I I P E Q V K G Q L T S S P R A Q G R R P Q S . E A P A G G . W I S S P N R . R D N . L A A P E P R D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S	L R A R S R A L Q T G P L P V A L P A W D P P A E S H . P A A S  GGGACGAAGGCCTCAAAGCTGAGAGGCCCCTGCCGGTGGTGATGGATATCATCCCCGAACAGGTGAAGGGACAACTGACTAGCAGCCCCAGAGCCC  GCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGGCCACCCAC
CAGGGACGAAGGCCTCAAAGCTGAGAGGCCCCTGCCGGTGGGTG	GGGACGAAGGCCTCAAAGCTGAGAGGCCCCTGCCGGTGGTGATGGATATCATCCCCGAACAGGTGAAGGGACCACTAGCAGCCCCAGAGCCC CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACCGCCCACTACCTATAGTAGGGGCTTGTCCACTTCCCTGTTGACTGATCGTCGGGGTCTCGGG  G T K A S K L R G P C R W V M D I I P E Q V K G Q L T S S P R A D G R R P Q S . E A P A G G . W I S S P N R . R D N . L A A P E P R D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S P
AGGGACGAAGGCCTCAAAGCTGAGAGGCCCCTGCCGGTGGGTG	GGGACGAAGGCCTCAAAGCTGAGAGGCCCCTGCCGGTGGTGATGGATATCATCCCCGAACAGGTGAAGGGACAACTGACTAGCAGCCCCAGAGCCC GCCTGCTTCCGGAGTTTCGACTCTCCGGGACCGCCCACCCCCTACCTA
G T K A S K L R G P C R W V M D I I P E Q V K G Q L T S S P R A CONTROL R R P Q S . E A P A G G . W I S S P N R . R D N . L A A P E P CR D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S	CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGCCACCCAC
CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGGCCACCCAC	CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGCCACCCAC
CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGCCACCCAC	CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGCCACCCAC
CCCTGCTTCCGGAGTTTCGACTCTCCGGGGACGGCCACCCAC	CCCTGCTTCCGGGGTTTCGACTCTCCGGGGACGGCCACCCAC
GTKASKLRGPCRWVMDIIPEQVKGQLTSSPRA GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQS	G T K A S K L R G P C R W V M D I I P E Q V K G Q L T S S P R A G R R P Q S . E A P A G G . W I S S P N R . R D N . L A A P E P R D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S P CCCTGCGGATCCCAGCCTAAAAGACCCCAGAGACCTCAGCTATGGAGCC
GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQS	GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQSP
GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQS	GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQSP
GRRPQS.EAPAGG.WISSPNR.RDN.LAAPEP RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQS	G R R P Q S . E A P A G G . W I S S P N R . R D N . L A A P E P R D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S P
RDEGLKAERPLPVGDGYHPRTGEGTTD.QPQS	R D E G L K A E R P L P V G D G Y H P R T G E G T T D . Q P Q S P
* <del>                                    </del>	CCCTGCGGATCCCAGCCTAAAAGACACCAGAGACCTCAGCTATGGAGCC
	CCCTGCGGATCCCAGCCTAAAAGACACCAGAGACCTCAGCTATGGAGCC
	ACCCTGCGGATCCCAGCCTAAAAGACACCAGAGACCTCAGCTATGGAGCC
CCCTGCGGATCCCAGCCTAAAAGACACCAGAGACCTCAGCTATGGAGCC	
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	TGGGACGCCTAGGGTCGGATTTTCTGTGGTCTCTGGAGTCGATACCTCGG

FIGURE 14C

3001	CTC	GCG	AGA	SCCG	GAC	CCG	GT	GTCG	AG	SCT	GCT(	CGA	CTP	TGC	CAAA	GGCC	SAAG	ACG	TCG	CCA	AGC	ACG	GCG	GCT	CG:	rgce	AG	GGG	3CG	TGC	TAG	AGT
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CCT	GCC	AGC	CTG	CTGC	GCG	CCG	GG	GCC	TG	CGG	rcg	CCT	ccc	CGGC	STC	CGG	CCGA	TCF	AGCC	CAGC	CCI	GTT	GCC	GGC	CC	ACTO	GC'	TAT	rga	GGC	CGI	CTC
GGA	CCGG	TCG	GAC	GAC	CGC	GGC	cc	CGG	AC	GCC <i>I</i>	AGC	GGA	GGC	ccc	CAGO	GCC	GCT	AGT	CGG	STCG	GGA	CAA	CGG	ccc	GG:	rgac	CG.	AT?	ACT	CCG	GCA	GAG
L	A	s	L	L	G	А	G	A	L	R	s	P	P	G	s	R	P	I	s	Q	P	С	С	R	P	т	R	Y	E	A		, s
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11C.	*****	WI.C	IGA	HCA.			- <del></del>	<del></del>		• • •	***	+++	++	++-	•••			•				+			•+	•••	-		•		+	<del></del> +
AAG	TACC	TAC	ACT	TGT	CGT	GAC	CT	CTT	GC.	ACC	TGG	TAG	AG	AGG	CGG	GGC	GGAC	GCC	CGAC	CAGA	CCC	CGAC	CTCC	CTAC	CTA	GAA	STT	CGZ	AAA	ACG	TGI	rgac
F	м	D	v	N :	s :	r V	Ą	R '	r	v i	D	н	L	s	A	T .	A (	: (	G (	C 1	. (	3.		3		s :	s	s	F	С	T	L
F	M	D +	v	N :	s :	r v	₽ • <del>  •</del>	R '	r <del>  • • •</del>	V 1	D	н	L	s	,A	т.	A (	: (	3 (	C I	. (	3 . <del>-  -</del> -	. (	3 •••	• • • •	s :	s • • •	s	F •   •	c	T	L · · · ·
• • • •	<del></del>			+				•••		• • •		<del>-   -</del>		<del></del>	• • •		<del></del>								•••	•••	•••	••	•   •	•••		
ACC	CATA	TG1	rcec	CCT	<del></del>	rggz	AAC	AGC	ccc	ACG	GGG	<del>·   ·</del>	CA	<del>·   ·</del>	GCT	AGGA	GCCI	CA	ACT(	CAAC	CAGO		SCTO	CAG	<del>···</del> l	TCA	<del>· i ·</del> sgc	:CG	<del>+ →</del> ATG	AGG	GAC	CAGA
ACC	CATA	TGT	cgc	CCT	ACC	rggz	· i ·	AGC	ccc	ACG	GGG	CCT	CA	CTA	GCT		GCCI	CA	ACT(	CAAC	CAGO	SAAC	SCTO	CAG	GCC	TCA	GGC	:CG	ATG	AGG	GAC	CAGA
ACC	CATA	TG1	CGC	CCT GGA	ACC:	rgg/	AAC +++ rtg	AGC	CCC I · · · GGG	ACG	GGG	CCT GGA	CA(	CTA GAT	GCT: CGA	AGGA	GCC1	CA	ACTO	CAAC	CAGO	SAAC	GAC	CAGG	GCC	TCA(	GGC	CG	ATG	AGG	GAC	CAGA

FIGURE 15

# INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/22604

IPC(7) US CL	SSIFICATION OF SUBJECT MATTER :Please See Extra Sheet. :Please See Extra Sheet. o International Patent Classification (IPC) or to both	national classification and IPC	
	DS SEARCHED		
	ocumentation searched (classification system follow	ed by classification symbols)	······································
<b>U.S</b> . :	Please See Extra Sheet.		
Documentat	ion searched other than minimum documentation to th	ne extent that such documents are included	in the fields searched
	ata base consulted during the international search (re Extra Sheet.	name of data base and, where practicable,	search terms used)
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
Α	US 5,739,307 A (JOHNSON, JR (14/04/98), see entire document.	. ET AL) 14 April 1998	1-38
A	US 5,747,655 A (JOHNSON, JR. ET see entire document.	AL) 05 May 1998 (05/05/98),	1-38
Furth	er documents are listed in the continuation of Box C	See patent family annex.	
*A* doc	cial categories of cited documents: ument defining the general state of the art which is not considered be of particular relevance	"T" later document published after the inter date and not in conflict with the applic the principle or theory underlying the	ation but cited to understand nvention
"L" doc	ier document published on or after the international filing date ument which may throw doubts on priority claim(s) or which is d to establish the publication date of another citation there is the restriction of the restric	"X" document of particular relevance; the considered novel or cannot be considered when the document is taken alone "Y" document of particular relevance; the	d to involve an inventive step
•	oial reason (as specified) ument referring to an oral disclosure, use, exhibition or other uns	considered to involve an inventive sembled with one or more other such being obvious to a person skilled in the	tep when the document is documents, such combination
	ument published prior to the international filing date but later than priority date claimed	"&" document member of the same patent f	amily
	actual completion of the international search MBER 1999	Date of mailing of the international search 10 FEB 2000	ch report
Commission Box PCT Washington,	ailing address of the ISA/US er of Patents and Trademarks D.C. 20231	Authorized officer  PREMA MERTZ  Telephone No. (703) 308-0196	2_

### INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/22604

# A. CLASSIFICATION OF SUBJECT MATTER: IPC (7):

C07K 14/47, 14/475; C12N 5/10, 15/12, 15/16, 15/63, 15/64; A61K 38/16, 38/17, 38/18, 39/395, 48/00

A. CLASSIFICATION OF SUBJECT MATTER:

US CL :

530/350,399; 536/23.1, 23.5, 23.51, 24.3, 24.31; 514/2, 8, 12, 44; 435/69.1, 69.4, 70.1, 71.1, 71.2, 471, 325, 375, 252.3, 254.11, 320.1, 6, 7.1, 7.2

#### B. FIELDS SEARCHED

Liminum documentation searched

Classification System: U.S.

530/350,399; 536/23.1, 23.5, 23.51, 24.3, 24.31; 514/2, 8, 12, 44, 45/69.1, 69.4, 70.1, 71.1, 71.2, 471, 325, 375, 252.3, 254.11, 320.1, 6, 7.1, 7.2

### B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

WEST, CAS ONLINE, MEDLINE, EMBASE, BIOSIS search terms: artemin, growth factor, polypeptide, polynucleotide, cDNA, GFR alpha, recombinant production, antibody, detection, assay, hybridization, treatment, method, therapy.

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- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

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(54) Title: ARTEMIN, A NOVEL NEUROTROPHIC FACTOR

(57) Abstract: A novel growth factor, artemin, which belongs to the GDNF/neurturin/persephin family of growth factors, is disclosed. The human and mouse amino sequences have been identified. Human and mouse artemin genomic DNA sequences have been cloned and sequenced and the respective cDNA sequences identified. In addition, methods for treating degenerative conditions using artemin, methods for detecting artemin gene alterations and methods for detecting and monitoring patient levels of artemin are provided.

# ARTEMIN, A NOVEL NEUROTROPHIC FACTOR

# Reference to Government Grant

This invention was made with government support under NIH Grant Number 5RO1-AG13730-03. The government has certain rights in this invention.

## 5 Related Applications

This application, which claims priority to Provisional Application Serial No. 60/108,148 filed November 12, 1998, is a continuation-in-part of Application Serial No. 09/163,283, filed September 29, 1998.

### 10 Background of the Invention

### (1) Field of the Invention

This invention relates generally to trophic or growth factors and, more particularly, to a new growth factor, artemin, which is a member of the neurturin-persephin-GDNF family of growth factors.

### 15 (2) Description of the Related Art

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The development and maintenance of tissues in complex organisms requires precise control over the processes of cell proliferation, differentiation, survival and function. A major mechanism whereby these processes are controlled is through the actions of polypeptides known as "growth factors". These structurally diverse molecules act through specific cell surface receptors to produce these actions.

Growth factors termed "neurotrophic factors" promote differentiation, maintain a mature phenotype and provide trophic support, promoting growth and survival of neurons. Neurotrophic factors reside in the nervous system or in innervated tissues. Nerve growth factor (NGF) was the first neurotrophic factor to be identified and characterized (Levi-Montalcini et al., J. Exp. Zool. 116:321, 1951). 5 NGF exists as a non-covalently bound homodimer that promotes the survival and growth of sympathetic, neural crest-derived sensory, and basal forebrain cholinergic neurons. In sympathetic neurons this substance produces neurite outgrowth in vitro and increased axonal and dendritic growth in vivo. (See Levi-Montalcini and Booker, Proc Nat'l Acad Sci 46:384-391, 1960; Johnson et al. Science 210: 916-918, 1980; 10 Crowley et al., Cell 76:1001-12, 1994). NGF has effects on cognition and neuronal plasticity, and can promote the survival of neurons that have suffered damage due to a variety of mechanical, chemical, viral, and immunological insults (Snider and Johnson, Ann Neurol 26:489-506, 1989; Hefti, J Neurobiol 25:1418-35, 1994). NGF also is known to extensively interact with the endocrine system and in immune and 15 inflammatory processes. (Reviewed in Scully and Otten, Cell Biol Int 19:459-469, 1995; Otten and Gadient, Int. J. Devl Neurosci 13:147-151, 1995). For example, NGF promotes the survival of mast cells. (Horigome et al. J Biol Chem 269:2695-2707, 1994).

In recent years it has become apparent that growth factors fall into classes, i.e. families or superfamilies based upon the similarities in their amino acid sequences. These families include, for example, the fibroblast growth factor family, the neurotrophin family and the transforming growth factor-beta (TGF-\$\beta\$) family. As an example of family member sequence similarities, TGF-\$\beta\$ family members have 7 canonical framework cysteine residues which identify members of this superfamily.

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NGF is the prototype of such a family of growth factors. Brain-derived neurotrophic factor (BDNF), the second member of this family to be discovered, was shown to be related to NGF by virtue of the conservation of all six cysteines that form the three internal disulfides of the NGF monomer (Barde, *Prog Growth Factor Res* 2:237-248, 1990 and Liebrock et al. *Nature 341*:149-152, 1989). By utilizing the information provided by BDNF of the highly conserved portions of two factors,

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additional members (NT-3, NT-4/5) of this neurotrophin family were rapidly found by several groups (Klein, *FASEB J 8*:738-44, 1994).

Recently, a new family of neurotrophic factors has been identified whose members are not structurally related to NGF and other neurotrophins but are structurally similar to TGF-β. As described in U.S. Patent No. 5,739,307 and copending application 08/931,858, the known members of this subfamily of the TGFß superfamily include glial cell line-derived neurotrophic factor (GDNF), neurturin (NTN), and persephin (PSP). The placement of GDNF, neurturin and persephin into the same growth factor family, also referred to as the GDNF ligand family, is based on the similarities of their physical structures and biological activities. Human persephin has about 40% sequence identity and about 43% sequence conservation with human GDNF; and about 49% sequence identity and about 50% sequence conservation with human neurturin. Similarly, human neurturin has about 43% sequence identity and about 53% sequence conservation with human GDNF. In addition, these three proteins have seven cysteine residues whose positions are exactly conserved. GDNF, neurturin and persephin each support the survival of dopaminergic midbrain neurons, and spinal and facial motor neurons, in both in vitro survival and in vivo injury paradigms, identifying these ligands as potential therapeutic agents in the treatment of neurodegenerative diseases (Henderson et al., Science 266, 1062-1064, 1994; Horger et al., J Neurosci. 18, 4929-37 1998; Lin et al., Science 260, 1130-1132, 1993; Milbrandt et al., Neuron 20, 245-53, 1998; Oppenheim et al., Nature 373, 344-346, 1995), reviewed by (Grondin and Gash, J Neurol. 245(11 Suppl 3), 35-42, 1998). However, while GDNF and neurturin both support the survival of many peripheral neurons in culture, including sympathetic, parasympathetic, sensory, and enteric neurons (Buj-Bello et al., Neuron 15, 821-828, 1995; Ebendal et al., J Neurosci Res 40, 276-284, 1995; Heuckeroth et al., Dev Biol 200, 116-29, 1998; Kotzbauer et al., Nature 384, 467-470, 1996; Trupp et al., J of Cell Biology 130, 137-148, 1995), persephin does not share any of these activities on peripheral neurons (Milbrandt et al., supra.

It was recently reported that GDNF and neurturin share receptors and signal transduction pathways (Creedon et al., *Proc. Natl. Acad. Sci. US 94*:7018-7023, 1997; Durbec et al., *Nature 381*:789-793, 1996; Trupp et al., *Nature 381*:785-789, 1996;

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Baloh et al., *Neuron 18*:793-802, 1997). These proteins act through a multicomponent receptor complex in which a transmembrane signal transducing component, the Ret protein-tyrosine kinase (Ret or Ret PTK), is activated upon the binding of a growth factor of the GDNF/neurturin family with a member of a family of closely related co-receptors named GFRα. A characteristic feature of the GFRα co-receptor family, is that its members have no transmembrane domain and are attached to the cell surface via a glycosyl-phosphatidylinositol (GPI) linkage (Durbec et al., *Nature 381*:789-793, 1996; Jing et al., *Cell 85*:1113-1124, 1996; Treanor et al., *Nature 382*:80-83, 1996; Trupp et al., *Nature 381*:785-789, 1996; Baloh et al., 1997, *supra*). The members of the GFRα family include GFRα1 (previously known as GDNFRα, TrnR1 and RetL1), GFRα2 (previously TrnR2, NTNRα and RetL2), GFRα3 (previously TrnR3) (GFRα Nomenclature Committee, *Neuron 19(3)*:485, 1997) and possibly GFRα4, a receptor currently only identified in the chicken (cGFRα4) (Enokido et al., *Current Biology 8*, 1019-1022, 1998).

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PCT/US99/22604

Results from in vitro experiments by multiple groups together indicate that 15 GFR $\alpha$ 1/RET is the preferred receptor for GDNF, and GFR $\alpha$ 2/RET is the preferred receptor for NTN, however cross-talk between the different receptors is possible (Baloh et al., 1997, supra; Jing et al., 1996, supra; Jing et al., J Biol. Chem. 272, 33111-33117, 1997; Klein et al., Nature 387, 717-721,1997; Sanicola et al., Proc. Natl. Acad. Sci., USA 94, 6238-6243,1997; Suvanto et al., Hum. Molec. Genet. 6, 20 1267-1273,1997; Treanor et al., 1996, supra). Recent analysis of GFRα1-deficient mice indicated that GFRa1 is the only physiologically relevant GDNF receptor in kidney organogenesis and enteric nervous system development (Cacalano et al., Neuron 21, 53-62,1998; Enomoto et al., Neuron 21, 317-324,1998). However, 25 GDNF-deficient mice have greater losses in peripheral ganglia than GFRα1-deficient mice, suggesting that GDNF can utilize additional other receptors to support survival of peripheral neurons, likely GFRα2/Ret (Cacalano et al., supra; Enomoto et al., supra). Persephin cannot signal through either the GFRα1/RET or GFRα2/RET receptor complexes (Milbrandt et al., supra), but a recent report indicated that persephin binds to cGFRa4 and likely also signals through RET (Enokido et al., 30 supra).

GFRα3 was first identified as an expressed sequence tag (EST) that is homologous to GFRa1 and GFRa2 (Baloh et al., Proc. Natl. Acad. Sci USA 95:5801-5806, 1998, supra; Jing et al., 1997, supra; Naveilhan et al., Proc Natl Acad Sci USA 95, 1295-300, 1998; Widenfalk et al., Eur. J. Neurosci. 10, 1508-1517, 1998; Worby et al., J Biol Chem 273, 3502-3508, 1998). However, analysis in transformed cells indicated that GFRa3 could not form a functional receptor with RET for any of the known GDNF family ligands, GDNF, neurturin or persephin (Baloh et al., 1998, supra;). GFRa3 expression is much more restricted than GFRa1 or GFRa2, with high level expression observed only in developing peripheral nerve and ganglia (Baloh et al., 1998, supra; Naveilhan et al., supra; Widenfalk et al., supra; Worby et al., 1998). Furthermore, one report demonstrated that in sensory neurons of the trigeminal ganglion, GFRa3 is expressed in a population of neurons distinct from GFR $\alpha$ 1 and GFR $\alpha$ 2, and likely overlaps with RET (Naveilhan et al., supra). Although the structural similarity, expression, and functional data together suggested that GFRa3 can interact with RET, the work reported herein was the first to demonstrate such an interaction and the first to identify a ligand for GFRa3.

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It is now generally believed that neurotrophic factors regulate many aspects of neuronal function, including survival and development in fetal life, and structural integrity and plasticity in adulthood. Since both acute nervous system injuries as well as chronic neurodegenerative diseases are characterized by structural damage and, possibly, by disease-induced apoptosis, it is likely that neurotrophic factors play some role in these afflictions. Indeed, a considerable body of evidence suggests that neurotrophic factors may be valuable therapeutic agents for treatment of these neurodegenerative conditions, which are perhaps the most socially and economically destructive diseases now afflicting our society. Nevertheless, because different neurotrophic factors can potentially act preferentially through different receptors and on different neuronal or non-neuronal cell types, there remains a continuing need for the identification of new members of neurotrophic factor families for use in the diagnosis and treatment of a variety of acute and chronic diseases of the nervous system.

PCT/US99/22604 WO 00/18799 6

# Summary of the Invention:

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Briefly, therefore, the present invention is directed to the new growth factor, artemin, which promotes cell survival and growth. Like GDNF and neurturin, artemin supports the survival of several peripheral neuron populations, and can also support dopaminergic neurons of the ventral midbrain. However, artemin is the only member of the GDNF family that binds to GFR $\alpha$  and activates the GFR $\alpha$ 3/RET receptor complex and, in addition, like GDNF and neurturin, artemin also binds to and activates GFRa1/RET. GFRa3 is described in patent application entitled "GFRa3, A Novel Member of the GDNF Coreceptor Family" filed concurrently with the present application on December 22, 1998 and this GFRa3 application is incorporated herein in its entirety by reference.

The present invention thus provides artemin polypeptides and polynucleotides. Artemin polypeptides within the scope of the present invention include any of the naturally occurring artemin polypeptides, conservatively substituted variants thereof or fragments thereof. The artemin polypeptides promote survival of peripheral and central neurons in culture, preferably in any one of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosinehydroxylase-expressing dopaminergic ventral midbrain neurons; more preferably, in any combination of these neurons and, most preferably, in all of these neurons.

Preferred artemin polypeptides include the predicted human mature polypeptides (SEQ ID NOS:3-5, see Figs. 3A, 3B, and 3C), and the predicted mouse mature polypeptides (SEQ ID NOS:34-36). These mature polypeptides can be generated by cleaving human or mouse pre-pro artemin (SEQ ID NOS: 26 and 29, respectively) or human or mouse pro-artemin (SEQ ID NOS:40 and 41, respectively), which are also within the scope of this invention, at one of the RXXR cleavage sites (see Figs. 1A, 1B, 1C and 2B). In addition, mature artemin can be generated by cleaving a polypeptide containing an N-terminal non-artemin pre-pro- region a mature artemin. Artemin polypeptides also include fragments from the first canonical cysteine to the seventh canonical cysteine of human or mouse mature artemin (SEQ ID NO:19 and SEQ ID NO:33, respectively).

Artemin is believed to show at least 75% sequence identity among homologous sequences from different mammalian species and, hence, mammalian

orthologs of mature artemin are believed to have at least 75% sequence identity with mature human artemin (SEQ ID NOS:3-5) and mature mouse artemin (SEQ ID NOS:34-36). Sequence homology may be as low as 65% in non-mammalian species such as avian species.

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Human artemin polypeptide is encoded by the cDNA contained in the clone deposited with the ATCC on December 22, 1998. Thus, human artemin polypeptide can be produced by transforming a host cell with the cDNA of this clone operably linked to expression regulatory elements. Conditions are then provided such that the cell expresses the encoded human artemin polypeptide. Such human artemin polypeptides are within the scope of the present invention.

The present invention also provides compositions comprising an artemin polypeptide and a pharmaceutically acceptable carrier suitable for administering to a cell to provide trophic support and/or to produce differentiation of that cell *in vitro* or  $ex\ vivo$  or to a cell in a patient. In order to facilitate the growth promoting effect of artemin, a GFR $\alpha$  coreceptor can be administered with artemin. Thus, in another embodiment, the present invention provides compositions which comprise an artemin polypeptide and a GFR $\alpha$  polypeptide such as GFR $\alpha$ 3 or GFR $\alpha$ 1.

In another embodiment, the present invention includes pan-growth factors and polynucleotides encoding the pan-growth factors. The pan-growth factors comprise a portion of the artemin polypeptide and a portion of at least one other growth factor from the TGF- $\beta$  superfamily. Preferably, the other growth factor can be neurturin, persephin or GDNF.

The present invention in another embodiment also provides isolated and purified and/or recombinant nucleic acid molecules comprising artemin nucleotide sequences. The artemin nucleotide sequences encode polypeptides consisting of artemin polypeptides, conservatively substituted variants thereof and fragments thereof. Also within the scope of this invention are fragments of artemin nucleotide sequences of at least 15 contiguous nucleotides. The artemin nucleic acid molecules specifically hybridize to mature human artemin nucleotide sequences (SEQ ID NOS:6-8) or their complements (SEQ ID NOS:9-11) or to mature mouse artemin nucleotide sequences (SEQ ID NOS:37-39) or their complements (SEQ ID NOS:60-62).

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Preferred polynucleotides of the present invention comprise the human polynucleotides which encode mature human artemin polypeptides (SEQ ID NOS:3-5), in particular, the polynucleotide sequences in SEQ ID NOS: 6-8 as depicted in Figs. 3A, 3B, and 3C or the corresponding mouse polynucleotides which encode mature mouse artemin polypeptides.(SEQ ID NOS:34-36), in particular, the polynucleotide sequences in SEQ ID NOS:37-39. Also encompassed by the present invention are human and mouse pro-artemin polynucleotides which encode proartemin polypeptides (SEQ ID NOS:40 and 41, respectively), in particular, the polynucleotide sequences in SEQ ID NOS:42 and 43, respectively, and human and mouse pre-pro artemin polynucleotides which encode pre-pro artemin polypeptides (SEQ ID NOS:26 and 29, respectively), in particular, the human polynucleotide sequences, SEO ID NOS:24, SEQ ID NO:30 or SEQ ID NO:46, and the mouse polynucleotide sequence, SEQ ID NO:27. In addition, polynucleotides can encode a polypeptide containing an N-terminal non-artemin pre-pro- region and a mature artemin such that upon expression of the polypeptide in an expression system and cleavage of the N-terminal non-artemin pre-pro region, a mature artemin is produced. Artemin polynucleotides also include portions of the artemin polynucleotide sequence encoding the polypeptide from the first canonical cysteine to the seventh canonical cysteine (SEQ ID NOS:19 and 33).

Sequences that are capable of hybridizing to human or mouse mature artemin polynucleotides can be used in methods for detecting the artemin gene and transcription products thereof, as well as in isolating artemin-encoding polynucleotides from other mammalian and non-mammalian species. Such methods are also within the scope of the present invention.

The present invention also provides antibodies which specifically react with artemin or a fragment thereof and methods for detecting artemin polypeptide in a sample by binding to the antibody.

In another embodiment, the present invention also provides methods for providing trophic support to and/or for producing differentiation of a target cell which comprises treating the cell with an effective amount of an artemin polypeptide. The target cell can be given artemin *in vitro*, *ex vivo* or *in vivo* in a patient suffering from a medical condition which causes degeneration or a loss of normal function of the target

cell. In one preferred embodiment, the cell is treated *in vivo* by administering to the patient an artemin polypeptide or an artemin polynucleotide encoding for expression the artemin polypeptide. The patient can be suffering from disease such as peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, nervous system tumors such as neuroblastomas, multiple sclerosis, infection, small cell lung carcinoma and enteric diseases such as idiopathic chronic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers.

Among the several advantages found to be achieved by the present invention, therefore, may be noted the provision of a new growth factor, artemin, in particular human artemin, which can be used to prevent the atrophy, degeneration or death of certain cells, in particular, neurons in need of trophic support; the provision of polynucleotides encoding artemin for use in gene therapy; the provision of methods for obtaining artemin by recombinant techniques; the provision of methods for providing trophic support to target cells, particularly neurons; the provision of methods for treating disease conditions involving cellular degeneration, and in particular, neuronal degeneration; the provision of methods that can detect and monitor artemin levels in a patient; and the provision of methods that can detect alterations in the artemin gene.

## Brief Description of the Drawings

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Figure 1A illustrates complementary human artemin genomic nucleotide sequences (SEQ ID NOS:1-2) which were assembled as described in the text, with the amino acids (SEQ ID NO:12) encoded by one of the reading frames shown below the nucleotide sequences and the possible RXXR proteolytic processing sites indicated by boxes.

Figure 1B illustrates the complementary cDNA sequences (SEQ ID NOS:24 and 25) and amino acid sequence (SEQ ID NO:26) of human pre-pro-artemin.

Figure 1C illustrates the complementary cDNA sequences (SEQ ID NOS:27 and 28) and amino acid sequence (SEQ ID NO:29) of mouse pre-pro-artemin.

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Figure 1D illustrates the complementary cDNA sequences (SEQ ID NOS:30 and 31) and amino acid sequence (SEQ ID NO:32) starting from a second methionine found in some human artemin cDNA clones which represent alternatively spliced human artemin mRNA;

Figure 2A illustrates an alignment of the amino acid sequences for the predicted mature forms of human GDNF (hGDNF, SEQ ID NO:13), human neurturin (hNTN, SEQ ID NO:14), human persephin (hPSP, SEQ ID NO:15) and human artemin (hART, SEQ ID NO:3) with identical amino acid residues enclosed in boxes and gaps inserted by the alignment program indicated by dashes;

Figure 2B illustrates the alignment of human and mouse pre-pro artemin (SEQ ID NOS:26 and 29, respectively) with the human and mouse putative signal sequence, pre- region, shaded (amino acids 1-39, SEQ ID NO:51 and SEQ ID NO:52, respectively), putative RXXR cleavage sites designated by thick lines, and arrowheads denoting the location of the two introns in the artemin gene.

Figure 2C is a schematic diagram of the location of the two introns in the artemin gene, and their splicing to produce artemin mRNA, with the location of a second starting methionine identified by RACE PCR in cDNA's isolated from some human cDNA libraries indicated by Met\*.

Figure 3A illustrates the amino acid sequence (SEQ ID NO:3) and the coding sequence (SEQ ID NO:6) of a first predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:9).

Figure 3B illustrates the amino acid sequence (SEQ ID NO:4) and the coding sequence (SEQ ID NO:7) of a second predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:10).

Figure 3C illustrates the amino acid sequence (SEQ ID NO:5) and the coding sequence (SEQ ID NO:8) of a third predicted mature human artemin polypeptide as well as the complement of the coding sequence (SEQ ID NO:11).

Figure 4 illustrates the family member sequence identity in the region between the first and seventh canonical framework cysteine residues aligned beginning with the first canonical framework cysteine for human GDNF (hGDNF, SEQ ID NO:16), human neurturin (hNTN, SEQ ID NO:17), human persephin (hPSP, SEQ ID NO:18) and human artemin (hART, SEQ ID NO:19).

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PCT/US99/22604 WO 00/18799 11

Figure 5 illustrates expression of artemin in human adult and fetal tissues showing in Fig. 5A an RNA spot blot containing poly(A) RNA's isolated from the human tissues shown in Fig. 5B and probed with a fragment of the human artemin cDNA.

Figures 6A and 6B illustrate in situ hybridization analysis of artemin expression in embryonic rat using a 32P-labeled RNA probe generated from a rat artemin cDNA fragment as a probe, showing in Fig. 6A a sagittal section of an embryonic day 14 (E14) rat, with the section oriented such that the top is the rostral and the left side is the dorsal, in which no signal is observed in the dorsal root ganglia (DRG) and strong signal is observed in probable exiting nerve roots (NR), with the arrowhead indicating expression observed in tissue below the liver that likely represents a lateral extension of the superior mesenteric artery, and showing in Fig. 6B a parasagittal section of E14 rat in the same orientation as in (A), in which high level artemin expression surrounding the developing superior mesenteric artery (SMA) is detected.

Figure 6C illustrates semi-quantitative RT-PCR analysis of artemin expression in Schwann cells isolated from neonatal rats (PS), from the adult sciatic nerve (N), and from the distal segment of the sciatic nerve at the indicated times following nerve transection.

Figure 7 illustrates that artemin supports the survival of peripheral and central neurons in culture, in which: Figs 7A and 7B show histograms of the number of surviving dorsal root ganglion (DRG) neurons cultured from post-natal day 1 (P1) rats in the presence of 50 ng/ml of the indicated factors (Fig. 7A, mean and SEM are shown, n=5-11) or in the presence of the indicated amounts of artemin (Fig. 7B, mean and SEM are shown, n=5); Fig. 7C shows a histogram of number of surviving trigeminal ganglion (TG) neurons cultured from P1 rats in the presence of 50 ng/ml of the indicated growth factor (mean and SEM are shown, n=4); Fig. 7D shows a histogram of the number of surviving nodose ganglion (NG) neurons cultured from P0 rats in the presence of BDNF (100 ng/ml) or the indicated GDNF family ligands (50 ng/ml) (mean and SEM from a representative experiment are shown, n=2); Fig. 7E shows a histogram of the number of surviving superior cervical ganglion (SCG) from P0 rats maintained in the presence of NGF for 5 days and then cultured in the

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presence of 50 ng/ml of the indicated factors (mean and SEM are shown, n=2-8); and Fig. 7F shows a histogram of the number of tyrosine hydroxylase-expressing (TH+) dopaminergic ventral midbrain neurons from E14 rats (EVM) cultured in the presence of 50 ng/ml of the indicated factors (mean and SEM are shown, n=9-15).

Figure 8 illustrates that artemin induces differentiation and proliferation in neuroblastoma cell lines showing in Figs. 8A-8C photographs of SK-SY5Y neuroblastoma cells cultured in the presence of no factor (Cn) (Fig. 8A), 10 μM all-trans retinoic acid (RA), a known inducer of differentiation of these cells (Fig. 8B), or 50 ng/mL artemin (ART) (Fig. 8C), and in Fig. 8D a histogram of BrdU incorporation by NBL-S neuroblastoma cells in the presence of 50 ng/mL of the indicated factor.

Figure 9A illustrates that artemin activates RET and downstream signaling in primary cultured SCG neurons showing an immunoblot of tyrosine phosphorylated RET or phosphorylated MAPK (MAPK) in lysates from SCG neurons treated with either no factor (Cn), GDNF, or artemin (ART) at 50 ng/ml.

Figure 9B illustrates that artemin activates RET in NBL-S neuroblastoma cells, showing an immunoblot of tyrosine phosphorylated RET or phosphorylated MAPK (MAPK) in lysates from NBL-S cells stimulated as in Fig. 9A in the absence or presence of the enzyme PI-PLC, which specifically cleaves GPI-anchored proteins from the cell surface.

Figures 9C-9E illustrate the direct binding of GFR $\alpha$ 1-Fc, GFR $\alpha$ 2-Fc, and GFR $\alpha$ 3-Fc receptor bodies to GDNF ligand family members showing graphs of the amount of absorbance at 450 nm plotted against increasing concentrations of the indicated soluble GFR $\alpha$ -Fc fusion protein added to microtiter plates coated with GDNF, neurturin, artemin or persephin, which were then treated with anti-human Fc antibodies conjugated to horseradish peroxidase (HRP) and binding of receptor bodies measured using the chromogenic HRP substrate 3,3',5,5'-tetramethylbenzidine, with error bars representing the standard deviation of duplicates from a representative experiment.

Figure 10 illustrates receptor activation by GDNF ligand family members in the presence of defined coreceptor components showing bar graphs of the amount of luciferase expression induced by the indicated growth factor in RET-expressing MG87 fibroblasts (RET-3T3) (Fig. 10A) or in NLF neuroblastoma cells (NLF) (Fig.

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10B) transiently transformed with the Gal4-Elk/Gal4-Luc reporter system together with an expression plasmid for the indicated coreceptor or the CMV plasmid with no insert (Control), with fold-activation determined by dividing luciferase activity in the indicated treatment condition by the no treatment control, and error bars representing the standard deviation of duplicate measurements from a representative experiment.

Figure 11 is a schematic diagram of ligand/receptor interactions in the GDNF ligand family deduced from multiple experimental paradigms *in vitro*, with large arrows indicating preferred ligand receptor interactions and smaller arrows indicating alternate receptor interactions.

Figure 12 illustrates the alignment of the amino acid sequences of human and mouse GFR $\alpha$ 3 precursor (SEQ ID NOS:63 and 64, respectively), with identical residues enclosed in boxes, the N-terminal signal sequence shaded (amino acids 1-31 in hGFR $\alpha$ 3 and 1-28 in mGFR $\alpha$ 3), mature GFR $\alpha$ 3 (amino acids 32-372 in hGFR $\alpha$ 3 and 29-369 in mGFR $\alpha$ 3), a shaded C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide shaded (amino acids 373-400 of hGFR $\alpha$ 3 and 370-397 of mGFR $\alpha$ 3) and putative N-linked glycosylation sites marked by black dots.

Figure 13 illustrates the human nucleotide sequence encoding GFRα3 precursor (SEQ ID NO:65) which includes the encoding sequence for the N-terminal signal sequence (nucleotides 1-93), the encoding sequence for mature GFRα3 (nucleotides 94-1116) and the encoding sequence for the C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide (nucleotides 1117-1203).

Figure 14A-C illustrate the complementary human genomic sequence that includes the artemin gene (SEQ ID NOS: 68 and 69) and the sequences of the three reading frames (SEQ ID NOS:70-72).

Figure 15 illustrates the partial rat artemin cDNA complementary sequences (SEQ ID NOS:73 and 74) and the encoded amino acid sequence (SEQ ID NO:75).

## Description of the Preferred Embodiments

The present invention is based upon the identification, isolation and sequencing of genomic and cDNA clones that encode a new member of the neurturin/persephin/ GDNF ligand family identified herein as artemin (referenced in copending application 09/163,283 as GF4). The discovery of artemin follows that of

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WO 00/18799 PCT/US99/22604

neurturin which is described in copending U.S. Application Serial No. 08/775,414, which is incorporated in its entirety by reference, and persephin which is described in copending U.S. Applications Serial Nos. 08/981,739 and 08/931,858 both of which are incorporated in their entirety by reference.

Artemin promotes the survival of several peripheral and central neuronal populations *in vitro*, including sympathetic neurons, neural crest, placodally-derived sensory neurons, and dopaminergic midbrain neurons. Artemin activates both the  $GFR\alpha 3/RET$  and  $GFR\alpha 1/RET$  receptor complexes *in vitro*, but it is believed  $GFR\alpha 3/RET$  is the preferred multicomponent receptor for artemin signaling *in vivo*.

As described in more detail below, artemin was discovered by using the murine pre-pro neurturin amino acid sequence to scan the High Throughput Genome Sequences (hgts) database. This search identified two hgts sequences (AC005038 and AC005051) which had regions of DNA homologous to a DNA encoding pre-pro neurturin. Although, the hgts sequences were later discovered to have numerous sequence errors, i.e. omissions, additions and incorrect bases, one of the hgts sequences (AC005038) had a stretch of 197 nucleotides (nucleotides 67,464 to 67,660) which ultimately turned out to be identical to nucleotides 663-467 of the complementary strand of the Artemin nucleic acid (Fig. 1B) and the other (AC005051) had a stretch of 183 nucleotides (nucleotides 113,379 to 113,561) identical to nucleotides 648-468 of the complementary strand of the Artemin nucleic acid (Fig. 1B).

In order to determine whether the hgts sequences corresponded to coding regions of a new growth factor, oligonucleotide primers were designed based upon the hgts sequences to amplify overlapping polynucleotides of the suspected coding region from human genomic DNA. The overlapping polynucleotides were sequenced and this sequence information was combined with the hgts sequences to assemble the complementary genomic nucleotide sequences (SEQ ID NO:1-2) shown in Fig. 1A. One of the reading frames in SEQ ID NO:1 encodes a partial pro-artemin amino acid sequence that extends from within a pro-domain including three potential RXXR proteolytic processing sites (boxes in Fig. 1A) and terminates at the second glycine residue shown in the last line of Figure 1A. Alignment of the predicted mature human artemin (hART) protein with the other GDNF ligands (Figure 2A) confirmed that

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artemin represents a new member of the GDNF ligand family, with artemin being most similar to neurturin and persephin (~45% identity), and somewhat more divergent from GDNF (~36% identity). Primers designed from the human artemin genomic sequence were used to identify a clone containing the mouse artemin gene.

To isolate full length mRNA species encoding artemin, primers corresponding to the human and mouse artemin genomic sequences were used to perform rapid amplification of cDNA ends (RACE) PCR on multiple human and mouse tissue cDNA libraries. Analysis of the full-length cDNAs from human and mouse (Figs 1B and 1C) enabled the prediction of full-length human and mouse artemin proteins (Figs. 1B, 1C and 2B) having a signal peptide for secretion (amino acids 1- 39 of Fig. 2B), and a large pro-region separated from the mature region by multiple conserved RXXR furin protease cleavage sites (Fig. 2B). Comparison of the human mRNA sequence with the genomic locus revealed the presence of two introns in the artemin coding region (Figure 2B and 2C), the second of which is positioned similarly to the introns found in the pro-domains of other GDNF family ligands.

Reference to mature artemin herein is intended to be construed to include growth factors of any origin which are substantially homologous to and which are biologically equivalent to mature human and mouse artemin characterized and described herein. Such substantially homologous growth factors may be native to any tissue or species and, similarly, biological activity can be characterized in any of a number of biological assay systems. Reference to pre-pro-artemin is intended to be construed to include pre-pro growth factors containing a pre- or leader or signal sequence region, a pro- sequence region and a mature artemin amino acid sequence as defined herein. Reference to pro-artemin is intended to mean a polypeptide lacking the signal sequence region, but containing both a pro-region ending in an RXXR cleavage sequence and a mature artemin amino acid sequence.

The terms "biologically equivalent" are intended to mean that the compositions of the present invention are capable of demonstrating some or all of the same growth promoting properties in a similar fashion, although not necessarily to the same degree as the recombinantly produced human artemin identified herein.

By "substantially homologous" it is meant that the degree of sequence identity between artemin orthologs including human, mouse and artemin from any other

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species, is greater than that between paralogs such as human artemin and human neurturin or human artemin and human persephin, and greater than that reported previously for members of the TGF-ß superfamily (for discussion of homology of TGF-ß superfamily members see Kingsley, *Genes and Dev 8*:133-46, 1994).

16

Sequence identity or percent identity is intended to mean the percentage of same residues between two sequences aligned using the Clustal method (Higgins et al, Cabios 8:189-191, 1992) of multiple sequence alignment in the Lasergene biocomputing software (DNASTAR, INC, Madison, WI). In this method, multiple alignments are carried out in a progressive manner, in which larger and larger alignment groups are assembled using similarity scores calculated from a series of pairwise alignments. Optimal sequence alignments are obtained by finding the maximum alignment score, which is the average of all scores between the separate residues in the alignment, determined from a residue weight table representing the probability of a given amino acid change occurring in two related proteins over a given evolutionary interval. Penalties for opening and lengthening gaps in the alignment contribute to the score. The default parameters used with this program are as follows: gap penalty for multiple alignment = 10; gap length penalty for multiple alignment = 10; k-tuple value in pairwise alignment = 1; gap penalty in pairwise alignment = 3; window value in pairwise alignment = 5; diagonals saved in pairwise alignment = 5. The residue weight table used for the alignment program is PAM250 (Dayhoff et al., in Atlas of Protein Sequence and Structure, Dayhoff, Ed., NBRF, Washington, Vol. 5, suppl. 3, p. 345, 1978).

To determine percent sequence identity between two sequences, the number of identical amino acids in the aligned sequences is divided by the total number of amino acids in the reference sequence. As used herein, the reference sequence is human artemin when determining its percent identity with mouse artemin and with the non-artemin growth factors, human neurturin, human GDNF or human persephin. Similarly, the reference sequence would be mouse artemin when determining its percent identity with mouse GDNF, mouse neurturin or mouse persephin and the reference sequence would be rat artemin when determining its percent identity with rat GDNF, rat neurturin and rat persephin. Referencing is to human neurturin when

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determining percent identity between human neurturin and non-human neurturin or between human neurturin and its human paralogs GDNF and persephin.

Percent conservation is calculated from the above alignment by adding the number of identical residues to the number of positions at which the two residues represent a conservative substitution (defined as having a log odds value of greater than or equal to 0.3 in the PAM250 residue weight table) and dividing by the total number of amino acids in the reference sequence. Preferred conservative amino acid changes are: R-K; E-D, Y-F, L-M; V-I, Q-H.

Table 1 shows the percent identity (I) and conservation (C) for comparisons of mature artemin, mature persephin, mature neurturin and mature GDNF from various species. Comparisons were made between mature human artemin (hART) and mature mouse artemin (mART) and between mature human artemin and mature human persephin (hPSP), mature human neurturin (hNTN) or mature human GDNF (hGDNF) using the alignments shown in Figs. 2A and 2B and the preferred conservative substitutions stated above. The persephin and neurturin comparisons were made as described in copending application Serial No. 08/931,858, with the first listed sequence being the reference sequence.

WO 00/18799 PCT/US99/22604

18
Table 1.

	COMPARISON	% IDENTITY	% CONSERVATION
	hART v. mART	88	90
	hART v. hPSP	45	48
5	hART v. hNTN	49	51
	hART v. hGDNF	36	40
	hPSP v. mPSP	81	81
	hPSP v. rPSP	80	81
10	mPSP v. rPSP	94	96
	hPSP v. hNTN	49	50
	hPSP v. hGDNF	40	43
	hNTN v. mNTN	90	93
15	hNTN v. rGDNF	44	53
	hNTN v. mGDNF	43	52
	hNTN v. hGDNF	43	53
	mNTN v. rGDNF	42	52
	mNTN v. mGDNF	41	51
20	mNTN v. hGDNF	41	52

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The sequence identity between human artemin and mouse artemin is about 88%. The persephin comparisons shown in Table 1 indicate that the identity between human persephin and mouse or rat persephin is about 80% whereas the degree of identity between mouse and rat persephin is about 94%. The neurturin comparisons in Table 1 indicate that mature mouse and human neurturin proteins have about 90% sequence identity. Furthermore, all artemin, persephin and neurturin orthologs of non-human mammalian species are believed to similarly have at least about 75% sequence identity with human artemin, human persephin, or human neurturin, respectively. For artemin, persephin or neurturin orthologs from non-mammalian species such as avian species, it is believed that the degree of homology with human artemin, human persephin, or human neurturin is at least about 65% identity.

By way of comparison, the variations between family members of the GDNF ligand family of growth factors can be seen by the comparisons shown in Table 1. For example, human artemin has about 49% sequence identity with human neurturin and about 36% sequence identity with human GDNF. Human persephin has about 49% sequence identity with human neurturin and about 40% sequence identity with human GDNF. Similarly, human neurturin has about 43% sequence identity with human GDNF. Thus, it is believed that any other member of the GDNF ligand family will have a similar sequence identity of about 40% of that of artemin, neurturin, persephin or GDNF of the same species and within a range of about 30% to about 60% sequence identity with artemin, neurturin, persephin or GDNF of the same species.

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Based on the data in Table 1, a given member of the GDNF ligand family would be expected to have lesser sequence identity with any other family member of the same species than the sequence identity present in orthologs of that family member in other species just as human GDNF and human neurturin are more closely related to mouse GDNF and mouse neurturin, respectively, than to each other or to GDNF (see U.S. Patent NO. 5,739,307). Similarly, any given member of the GDNF ligand family would be expected to have greater sequence identity with another family member than to any other known member of the TGF- $\beta$  superfamily (Kingsley, *supra*).

The conservation between members of the GDNF ligand family is also seen in Figure 4, which shows an alignment, using the Clustal program described above, of the human sequences for GDNF, neurturin (NTN), persephin (PSP) and artemin (ART) from the first to the seventh framework cysteine residues. From this alignment, it is evident that artemin is closely related to the other family members in that artemin and persephin share 46 out of 96 residues (48%), artemin shares 48 out of 96 residues (50%) with neurturin, and artemin and GDNF share 36 out of 96 residues (38%).

Orthologs of pre-pro artemin in non-human mammalian species can be identified by virtue of the mature portion of the amino acid sequence having at least about 75% sequence identity with one of the predicted mature human artemin sequences disclosed herein and nonmammalian orthologs of pre-pro artemin can be

identified by virtue of the mature portion of the amino acid sequence having at least about 65% identity with a mature human artemin sequence.

Also included within the meaning of substantially homologous is any artemin polypeptide which may be isolated by virtue of cross-reactivity with antibodies specific to human artemin or whose encoding nucleotide sequences, including genomic DNA, mRNA or cDNA may be isolated through hybridization with the complementary sequences shown in Figs 1-3 or fragments thereof. It will also be appreciated by one skilled in the art that degenerate DNA sequences can encode human artemin and these are also intended to be included within the present invention.

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Conservatively substituted artemin proteins retaining the biological activity of naturally occurring artemin are also within the scope of the present invention. Conservative amino acid substitutions refer to the interchangeability of residues having similar side chains. Conservatively substituted amino acids can be grouped according to the chemical properties of their side chains. For example, one grouping of amino acids includes those amino acids have neutral and hydrophobic side chains (A, V, L, I, P, W, F, and M); another grouping is those amino acids having neutral and polar side chains (G, S, T, Y, C, N, and Q); another grouping is those amino acids having basic side chains (K, R, and H); another grouping is those amino acids having acidic side chains (D and E); another grouping is those amino acids having aliphatic side chains (G, A, V, L, and I); another grouping is those amino acids having aliphatic-hydroxyl side chains (S and T); another grouping is those amino acids having amine-containing side chains (N, Q, K, R, and H); another grouping is those amino acids having aromatic side chains (F, Y, and W); and another grouping is those amino acids having sulfur-containing side chains (C and M). Preferred conservative amino acid substitutions groups are: R-K; E-D, Y-F, L-M; V-I, and Q-H.

As used herein, an artemin polypeptide can also include modifications of the artemin sequences identified herein, including sequences in which one or more amino acids have been inserted, deleted or replaced with a different amino acid or a modified or unusual amino acid, as well as modifications such as glycosylation or phosphorylation of one or more amino acids so long as the polypeptide containing the modified sequence retains the biological activity of artemin. Inserted or deleted

amino acid(s) can be added to or removed from the N-terminus, C-terminus or within the naturally-occurring amino acid sequence. By retaining the biological activity, it is meant that the modified polypeptide can bind to and activate  $GFR\alpha 1/RET$  and/or  $GFR\alpha 3/RET$  expressed by a cell, although not necessarily at the same level of potency as that of the mature human artemin polypeptide identified herein. Assays for testing such activation are known in the art and include the Gal4-Elk/Gal4-Luc reporter system described in Example 7 below. The term "trophic support" is used herein to mean that a growth factor such as artemin provides nourishment to a cell such that the cell maintains or recovers at least one of its normal functions.

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It is intended that the term artemin polypeptide also includes naturally occurring allelic variants of the human artemin sequences disclosed herein. For example, cDNAs encoding a second starting methionine were identified by RACE PCR from some human cDNA libraries and would encode the pre-pro-artemin shown in Fig. 1D (SEQ ID NO:32). However, mature artemin generated from this variant would be identical to the mature human artemin shown in Fig. 3A (SEQ ID NO:3).

Fragments of artemin are also encompassed by the present invention. Such fragments may be of any length but preferably retain the biological activity of artemin or are antigenic. The minimum length of such biologically active or antigenic fragments can readily be determined by those skilled in the art using known techniques. Preferably, the minimum length of fragments of artemin is at least 8 amino acids, more preferably, at least 10 amino acids, still more preferably at least 12 amino acids, even still more preferably at least 15 amino acids and most preferably, at least 20 amino acids or greater. One fragment of artemin which is believed to retain the survival promoting activity of artemin begins at the first conserved cysteine residue and ends at the seventh conserved cysteine residue (Fig. 4, SEQ ID NO:19 (human) or SEO ID NO:33 (mouse)). Antigenic fragments are capable of eliciting artemin-specific antibodies when administered to a host animal and includes those smaller fragments that require conjugation to a carrier molecule to be immunogenic. Typically, antigenic fragments will be at least 5 or 6 amino acids in length and may be any length up to the length of mature artemin, preferably an antigenic fragment is 8, more preferably 10, still more preferably 12 amino acids in length, even still more

preferably 15 amino acids in length and most preferably 20 amino acids or more in length.

22

It is also believed that particular discrete fragments of artemin, or analogues thereof, can serve as an agonist where the fragment activates an artemin receptor, such as  $GFR\alpha 3/RET$  or  $GFR\alpha 1/RET$ , to elicit the survival or growth promoting action on a target cell, or other artemin fragments or analogues can serve as antagonists to artemin where they bind to, but do not activate, the receptor or do not promote survival and growth. Such fragments or analogues that are agonists and those that are antagonists are also within the scope of the present invention.

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Although it is not intended that the inventors herein be bound by any theory, it is thought that the human artemin protein identified herein as well as orthologs from other tissues and species may exist as dimers in their biologically active form in a manner consistent with what is known for other factors of the TGF-ß superfamily.

In addition to homodimers, the monomeric units of artemin dimers can be used to construct stable growth factor heterodimers or heteromultimers comprising at least one monomer unit of artemin. This can be done by dissociating a homodimer of artemin into its component monomeric units and reassociating in the presence of a monomeric unit of a second or subsequent homodimeric growth factor. This second or subsequent homodimeric growth factor can be selected from a variety of growth factors including neurturin, GDNF, persephin, a member of the NGF family such as NGF, BDNF, NT-3 and NT-4/5, a member of the TGF-ß superfamily, a vascular endothelial growth factor, a member of the CNTF/LIF family and the like. By forming heterodimers or heteromultimers of artemin and one or more other growth factors, the resultant hybrid growth factor would be expected to be able to bind to at least two distinct receptor types preferentially having a different tissue distribution. The resultant heterodimers or heteromultimers would be expected to show a different and, possibly, an enlarged spectrum of cells upon which it could act or to provide greater potency. It is also possible that the heterodimer or heteromultimer might provide synergistic effects not seen with homodimers or homomultimers. For example, the combination of factors from different classes has been shown to promote long-term survival of oligodendrocytes whereas single factors or combinations of

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factors within the same class promoted short-term survival (Barres et al., *Development 118*:283-295, 1993).

Heterodimers can be formed by a number of methods. For example, homodimers can be mixed and subjected to conditions in which dissociation/unfolding occurs, such as in the presence of a dissociation/unfolding agent, followed by subjection to conditions which allow monomer reassociation and formation of heterodimers. Dissociation/unfolding agents include any agent known to promote the dissociation of proteins. Such agents include, but are not limited to, guanidine hydrochloride, urea, potassium thiocyanate, pH lowering agents such as buffered HCl solutions, and polar, water miscible organic solvents such as acetonitrile or alcohols such as propanol or isopropanol. In addition, for homodimers linked covalently by disulfide bonds as is the case with TGF-β family members, reducing agents such as dithiothreitol and β-mercaptoethanol can be used for dissociation/unfolding and for reassociation/refolding.

Heterodimers can also be made by transforming a cell with two or more factors such that the transformed cell produces heterodimers as has been done with the neurotrophins. (Heymach and Schooter, *J Biol Chem 270*:12297-12304, 1995). Another method of forming heterodimers is by combining artemin homodimers and a homodimer from a second growth factor and incubating the mixture at 37°C.

When heterodimers are produced from homodimers, the heterodimers may then be separated from homodimers using methods available to those skilled in the art such as, for example, by elution from preparative, non-denaturing polyacrylamide gels. Alternatively, heterodimers may be purified using high pressure cation exchange chromatography such as with a Mono S cation exchange column or by sequential immunoaffinity columns.

It is well known in the art that many proteins are synthesized within a cell with a signal sequence at the N-terminus of the mature protein sequence and the protein carrying such a leader sequence is referred to as a preprotein. The pre- portion of the protein is cleaved during cellular processing of the protein. In addition to a pre-leader sequence, many proteins contain a distinct pro sequence that describes a region on a protein that is a stable precursor of the mature protein. Proteins synthesized with both pre- and pro- regions are referred to as preproproteins. In view of the processing

events known to occur with other TGF-ß family members, the inventors believe that the form of the artemin protein as synthesized within a cell is the pre-pro-artemin.

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Human and mouse pre-pro-artemin polypeptides are believed to preferably contain an N-terminal methionine of a signal sequence (pre-region) of 39 amino acids (Fig. 2B; SEQ ID NOS:48 and 49, respectively; amino acids 1-39 of SEQ ID NOS:26 and 29, respectively). It is known that the full length of a leader sequence is not necessarily required for the sequence to act as a signal sequence and, therefore, within the definition of pre-region of artemin is included fragments thereof, usually Nterminal fragments, that retain the property of being able to act as a signal sequence, that is to facilitate co-translational insertion into the membrane of one or more cellular organelles such as endoplasmic reticulum, mitochondria, golgi, plasma membrane and the like. It is also possible that the pre-pro-artemin polypeptide might have one or more isoforms having an N-terminal Leucine, inasmuch as isoforms of another growth factor, human fibroblast growth factor-2, have alternative initiations of translation at CUG start codons in addition to AUG (Arnaud et al., Molecular and Cellular Biology 19:505-514, 1999). Thus, isoforms of pre-pro-artemin could begin, for example, before the methionine (the first amino acid shown in Figure 1B) with the start point being at the CTG codon beginning at nucleotide 284 in the genomic human artemin sequence (see Figure 14) or after the same methionine with the start point being the CTG codon beginning at nucleotide 329 so long as the pre-region of the isoform can serve as a signal sequence. All such isoforms are within the scope of the present invention.

Moreover, also within the scope of the present invention are polypeptides containing a non-artemin pre-pro- region and a mature artemin as well as polynucleotides encoding such polypeptides. The polypeptides can generate a mature artemin upon cleavage of the non-artemin pre-pro- region and the polynucleotides can be used in an expression system to produce the polypeptide which upon cleavage of the non-artemin pre-pro- region yields mature artemin. Such non-artemin pre-pro- region polypeptides and encoding polynucleotides are well known in the art and, preferably, include a pre-pro- region from another growth factor such as neurturin or persephin although any pre-pro- region can be used so long as it functions as a pre-pro- region as described herein.

The artemin pre- region is followed by a pro- domain which is believed to preferably terminate with an RXXR consensus site immediately before the N-terminal amino acid of mature artemin. Thus, mature human and mouse artemin are believed to be preferably generated by proteolytic cleavage of human and mouse pro-artemin (SEQ ID NOS:40 and 41, respectively) after one of the three RXXR consensus sequences shown in Figs. 1A and 2B, (RARR, RGGR and RAAR for human and RTLR, RGAR and RAAR for mouse) although it is possible that cleavage could occur at some other, non-consensus site to produce additional isoforms. All such isoforms are included within the scope of the present invention.

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Cleavage after the first RXXR sequence following the N-terminus of human pro-artemin will generate the mature polypeptide shown in Fig. 3C (SEQ ID NO:5). Similarly, cleavage after the second and third RXXR sequences will generate the mature human polypeptides shown in Figs. 3B (SEQ ID NO:4) and 3A (SEQ ID NO:3), respectively. By analogy, predicted mature mouse artemin sequences are SEQ ID NOS:34, 35 and 36. Based on the biological assays described below, the preferred mature human artemin consists of SEQ ID NO:3 and similarly, the preferred mature mouse artemin consists of SEQ ID NO:34, respectively. The preferred human proregion polypeptide would thus comprise amino acids 40-107 of Fig. 2B (SEQ ID NO:50) and the corresponding preferred mouse pro-region polypeptide would comprise amino acids 40-111 of Fig. 2B (SEQ ID NO:51). Similarly the preferred human pre-pro- region polypeptide would thus comprise amino acids 1-107 of Fig. 2B (SEQ ID NO:52) and the corresponding preferred mouse pre-pro- region polypeptide would comprise amino acids 1-111 of Fig. 2B (SEQ ID NO:53). The mature, secreted artemin molecule is likely to form a disulfide linked homodimer by analogy to other members of the TGF-ß family.

The human and mouse preferred pre- region polynucleotides comprise nucleotides 1-117 of Fig. 1B (SEQ ID NO:54) and nucleotides 1-117 of Fig. 1C (SEQ ID NO:55), respectively; the human and mouse pro- region polynucleotides comprise nucleotides 118-321 of Fig. 1B (SEQ ID NO:56) and nucleotides 118-333 of Fig. 1C (SEQ ID NO:57), respectively; and the human and mouse pre-pro-region polynucleotides comprise nucleotides 1-321 of Fig. 1B (SEQ ID NO:58) and nucleotides 1-333 of Fig. 1C (SEQ ID NO:59), respectively. It is noted, however, that

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alternative start points and alternative non-consensus cleavage points as discussed above would result in coding sequences corresponding to the alternative isoforms of pre-pro-artemin and mature artemin.

A preferred artemin according to the present invention is prepared by recombinant DNA technology although it is believed that artemin can be isolated in purified form from cell-conditioned medium as was done for neurturin.

By "pure form" or "purified form" or "substantially purified form" it is meant that an artemin composition is substantially free of other proteins which are not artemin. Preferably, a substantially purified artemin composition comprises at least about 50 percent artemin on a molar basis compared to total proteins or other macromolecular species present. More preferably, a substantially purified artemin composition will comprise at least about 80 to about 90 mole percent of the total protein or other macromolecular species present and still more preferably, at least about 95 mole percent or greater.

Recombinant artemin may be made by expressing the DNA sequences encoding artemin in a suitable transformed host cell. Using methods well known in the art, the DNA encoding artemin may be linked to an expression vector, transformed into a host cell and conditions established that are suitable for expression of artemin by the transformed cell.

Any suitable expression vector may be employed to produce recombinant artemin such as, for example, the mammalian expression vector pCB6 (Brewer, *Meth Cell Biol 43*:233-245, 1994) or the *E. coli* pET expression vectors, specifically, pET-30a (Studier et al., *Methods Enzymol 185*:60-89, 1990). Other suitable expression vectors for expression in mammalian and bacterial cells are known in the art as are expression vectors for use in yeast or insect cells. Baculovirus expression systems can also be employed.

A number of cell types may be suitable as host cells for expression of recombinant artemin. Mammalian host cells include, but are not limited to, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo 205 cells, 3T3 cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from *in vitro* culture of primary tissue, primary explants, HeLa cells, mouse L cells, BHK, HL-60, U937,

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HaK and Jurkat cells. Yeast strains that may act as suitable host cells include Saccharomyces cerevisiae, Schizosaccharomyces pombe, Kluyveromyces strains, Candida, and any other yeast strain capable of expressing heterologous proteins. Host bacterial strains include Escherichia coli, Bacillus subtilis, Salmonella typhimurium and any other bacterial strain capable of expressing heterologous proteins. If the polypeptide is made in yeast or bacteria, it may be necessary to modify the polypeptide, for example, by phosphorylation or glycosylation of the appropriate sites using known chemical or enzymatic methods, to obtain a biologically active polypeptide.

27

PCT/US99/22604

The polypeptide of the invention can also be expressed in transgenic plants (see, for example, U.S. Patent No. 5,679,880) or transgenic animals such as, for example, cows, goats, pigs, or sheep whose somatic or germ cells contain a nucleotide sequence encoding human artemin.

The expressed artemin polypeptide can be purified using known purification procedures, such as gel filtration and ion exchange chromatography. Purification may also include affinity chromatography using an agent that will specifically bind the artemin polypeptide, such as a polyclonal or monoclonal antibody raised against a mature artemin or fragment thereof. Other affinity resins typically used in protein purification may also be used such as concanavalin A-agarose, heparin-toyopearl® or Cibacrom blue 3GA Sepharose®. Purification of artemin can also include one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether.

It is also contemplated that an artemin polypeptide may be expressed as a fusion protein to facilitate purification. Such fusion proteins, for example, include an artemin amino acid sequence fused to a histidine tag such as when expressed in the pET bacterial expression system as well as the artemin amino acid sequence fused to the amino acid sequence of maltose binding protein (MBP), glutathione-S-transferase (GST) or thioredoxin (TRX). Similarly, the polypeptide of the invention can be tagged with a heterologous epitope and subsequently purified by immunoaffinity chromatography using an antibody that specifically binds such epitope. Kits for expression and purification of such fusion proteins and tagged proteins are commercially available.

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Artemin and fragments thereof may also be produced by chemical synthesis using methods known to those skilled in the art.

Artemin may be expressed in the monomeric units or such monomeric form may be produced by preparation under reducing conditions. In such instances refolding and renaturation can be accomplished using one of the agents noted above that is known to promote dissociation/association of proteins. For example, the monomeric form can be incubated with dithiothreitol followed by incubation with oxidized glutathione disodium salt followed by incubation with a buffer containing a refolding agent such as urea.

Because multiple RXXR cleavage sites are present in the pro-artemin sequence, it is believed that isoforms of mature artemin exist. In addition, alternative non-consensus cleavage sites might also result in different isoforms. Thus, the mature artemin polypeptide may have a variable number of amino acids preceding the first canonical cysteine. Such alternate cleavage sites could be utilized differently among different organisms and among different tissues of the same organism. The N-terminal amino acids preceding the first of the seven conserved cysteines in the mature forms of members of the TGF-ß family vary greatly in both length and sequence. Furthermore, insertion of a ten amino acid sequence two residues upstream of the first conserved cysteine does not affect the known biological activities of one family member, dorsalin (Basler et al., *Cell 73*:687-702, 1993). By analogy, it is believed that artemin proteins containing sequences of different lengths preceding the first canonical cysteine may exist or could be made and that these would retain their biological activity.

The present invention also encompasses isolated polynucleotides comprising nucleotide sequences that encode any of the human and mouse artemin polypeptides described herein. As used herein, a polynucleotide includes DNA and/or RNA and thus the nucleotide sequences recited in the Sequence Listing as DNA sequences also include the identical RNA sequences with uracil substituted for thymine residues. Nucleotide sequences included in the invention are those encoding the human and mouse pre-pro-artemin amino acid sequences shown in Figs. 1B and 1C, respectively, as well as nucleotide sequences encoding the variant human artemin protein shown in Fig. 1D and nucleotide sequences encoding human and mouse pro-artemin (SEQ ID

NOS:40 and 41, respectively). Preferred polynucleotides which encode human prepro- and pro- artemin comprise SEQ ID NOS:24 and 42, respectively, and preferred polynucleotides encoding mouse pre-pro- and pro-artemin comprise SEQ ID NOS:27 and 43, respectively. Polynucleotides within the scope of this invention do not include isolated chromosomes.

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The invention also includes polynucleotides comprising a nucleotide sequence that encodes a mature artemin polypeptide such as mature human and mouse polypeptides consisting of any of SEQ ID NOS:3-5and SEQ ID NOS:34-36, respectively. Such polynucleotides encoding mature artemin include the human nucleotide sequences set forth in SEQ ID NOS:6-8 and the mouse nucleotide sequences set forth in SEQ ID NOS:37-39. A particularly preferred polynucleotide encodes SEQ ID NO:3 and comprises SEQ ID NO:6.

It is understood by the skilled artisan that degenerate nucleotide sequences can encode the artemin amino acid sequences described herein and these are also intended to be included within the present invention. For example, SEQ ID NO:44 represents the artemin coding sequence found in one human cDNA clone isolated by the inventors herein which contains G rather than A at position 582, but which does not change the encoded amino acid sequence. Such degenerate nucleotide sequences include modifications of naturally-occurring sequences in which at least one codon is substituted with a corresponding redundant codon preferred by a given host cell, such as *E. coli* or insect cells, so as to improve expression of recombinant artemin therein.

The present invention also encompasses vectors comprising an expression regulatory element operably linked to any of the artemin-encoding nucleotide sequences included within the scope of the invention. This invention also includes host cells, of any variety, that have been transformed with such vectors.

In yet another embodiment, a polynucleotide which specifically hybridizes to a human artemin-encoding polynucleotide or to its complement is provided. Specific hybridization is defined herein as the formation of hybrids between a polynucleotide, including oligonucleotides, and a specific reference polynucleotide (e.g., a polynucleotide comprising a nucleotide sequence complementary to a nucleotide sequence encoding human artemin) wherein the polynucleotide preferentially hybridizes to the specific reference polynucleotide over other non-artemin

polynucleotides. Specifically hybridizing oligonucleotides are typically at least 15 nucleotides in length and are preferably at least 17 to at least 20 nucleotides long. Other preferred lengths include at least 22 to at least 25 nucleotides. A polynucleotide that specifically hybridizes to a reference sequence can be of any length, for example, about 15 nucleotides up to about 100 nucleotides or up to about 1000 nucleotides or up to about 10,000 nucleotides or even greater. Specific hybridization is preferably done under high stringency conditions which, as well understood by those skilled in the art, can readily be determined by adjusting several factors during hybridization and during the washing procedure, including temperature, ionic strength, length of hybridization or washing times, and concentration of formamide (see for example, Sambrook, Fritsch and Maniatis., *Molecular Cloning: a Laboratory Manual, 2d Ed., Vols. 1-3,* Cold Spring Harbor Laboratory Press, Plainview N.Y. 11803, 1989)

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The present invention also includes nucleic acid sequences which encode for artemin polypeptides that have survival or growth promoting activity and that preferentially bind anti-human or anti-mouse artemin antibodies over other antibodies that do not bind to human or mouse artemin.

Methods are also provided herein for producing artemin. Preparation can be by isolation from conditioned medium from a variety of cell types so long as the cell type produces artemin. A second and preferred method involves utilization of recombinant methods by isolating a nucleic acid sequence encoding artemin, cloning the sequence along with appropriate regulatory sequences into suitable vectors and cell types, and expressing the sequence to produce artemin.

On the basis of the structural similarities of artemin to the sequences of neurturin, GDNF and persephin, artemin would be expected to promote the survival and growth of neuronal as well as non-neuronal cells. As discussed above, GDNF, neurturin and persephin influence a broad spectrum of neuronal populations in the peripheral and central nervous systems. Moreover, all other growth factors isolated to date have been shown to act on many different cell types (for example see Scully and Otten, *Cell Biol Int 19*:459-469, 1005; Hefti, *Neurotrophic Factor Therapy 25*:1418-1435, 1994 which are incorporated by reference). As an example of the actions of neurotrophic factors on non-neuronal tissues, the prototypical neurotrophic factor,

NGF, also acts upon mast cells to increase their number when injected into newborn rats (Aloe, *J Neuroimmunol 18*:1-12, 1988). In addition, mast cells express the trk receptor and respond to NGF such that NGF is a mast cell secretogogue and survival promoting factor (Horigome et al., *J Biol Chem 269*:2695-2707, 1994). Moreover, members of the TGF-ß superfamily act on many cell types of different function and embryologic origin.

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Thus, it is likely that artemin will have trophic activity on a variety of different neuronal cells, both peripheral and central, as well as on non-neuronal cells. With respect to neuronal cells, artemin supports the survival of neurons from all peripheral ganglia examined thus far, including sympathetic neurons and neural crest and placodally-derived sensory neurons, and also supports the survival of at least one population of CNS neurons, i.e., dopaminergic midbrain neurons. In addition, the inventors herein have detected artemin expression in a number of adult and fetal tissues, including heart, kidney, lung, peripheral leukocytes and bone marrow, which further supports the conclusion that artemin can provide trophic support to a variety of neuronal and non-neuronal cells. This suggests a role for artemin in hematopoiesis, inflammation, allergy, and cardiomyopathy. Artemin's trophic activity on any particular target cell type can be determined by routine experimentation using standard reference models.

The invention also contemplates synthetic, pan-growth factors comprising at least one active domain of artemin combined with at least one active domain of one or more other non-artemin growth factors. (For example, see Ilag et al., *Proc Nat'l Acad Sci 92*:607-611, 1995). These pan-growth factors would be expected to have the combined activities or other advantageous properties of artemin and the one or more other growth factors. As such, these pan-growth factors are believed to be potent and multispecific growth factors that are useful in the treatment of a wide spectrum of degenerative diseases and conditions including conditions that can be treated by any or all of the parent factors from which the active domains were obtained. Such pan-growth factors might also provide synergistic effects beyond the activities of the parent factors (Barres et al., *supra*).

Pan-growth factors within the scope of the present invention can include chimeric or hybrid polypeptides that are constructed from portions of fragments of at

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least two growth factors. Growth factors of the TGF-ß superfamily are structurally related having highly conserved sequence landmarks whereby family members are identified. In particular, seven canonical framework cysteine residues are nearly invariant in members of the superfamily (Kingsley, Genes & Dev 8:133-146, 1994 which is incorporated by reference). Chimeric polypeptide molecules can, therefore, be constructed from a sequence that is substantially identical to a portion of the artemin molecule, up to one or more crossover points, and one or more sequences each of which is substantially identical with a portion of another TGF-B superfamily member extending on the other side of the corresponding one or more crossover points. For example, a portion of the amino terminal end of the artemin polypeptide can be combined with a portion of the carboxy terminal end of a neurturin polypeptide or alternatively a portion of the amino terminal end of a neurturin polypeptide can be combined with a portion of the carboxy terminal end of an artemin polypeptide. Such portions of artemin or neurturin polypeptides are preferably from about 5 to about 95, more preferably from about 10 to about 90, still more preferably from about 20 to about 80 and most preferably from about 30 to about 70 contiguous amino acids and such portions of another, non-artemin or, as the case may be, non-neurturin TGF-B superfamily member are preferably from about 5 to about 95, more preferably from about 10 to about 90, still more preferably from about 20 to about 80 and most preferably from about 30 to about 70 contiguous amino acids. For example, a particular crossover point might be between the third and fourth canonical framework cysteine residues. The particular non-artemin TGF-\( \beta \) family member could be selected from family members including but not limited to transforming growth factor-\(\beta\)1 (TGF\(\beta\)1), transforming growth factor-\(\beta\)2 (TGF\(\beta\)2), transforming growth factor-B3 (TGFB3), inhibin B A (INHBA), inhibin B B (INHBB), the nodal gene (NODAL), bone morphogenetic proteins 2 and 4 (BMP2 and BMP4), the *Drosophila* decapentaplegic gene (dpp), bone morphogenetic proteins 5-8 (BMP5, BMP6, BMP7 and BMP8), the Drosophila 60A gene family (60A), bone morphogenetic protein 3 (BMP3), the Vg1 gene, growth differentiation factors 1 and 3 (GDF1 and GDF3), dorsalin (drsln), inhibin  $\alpha$  (INH $\alpha$ ), the MIS gene (MIS), growth factor 9 (GDF-9), glial-derived neurotrophic growth factor (GDNF), neurturin (NTN) and persephin. Furthermore, additional crossover points can be used to incorporate any desired

WO 00/18799 PCT/US99/22604

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number of artemin portions or fragments with portions or fragments of any one or more other family members.

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In constructing a particular chimeric molecule, the portions of artemin and portions of the other, non-artemin growth factor are amplified using PCR, mixed and used as template for a PCR reaction using the forward primer from one and the reverse primer from the other of the two component portions of the chimeric molecule. Thus, for example a forward and reverse primers are selected to amplify the portion of artemin from the beginning to the selected crossover point between the third and fourth canonical cysteine residues using a artemin-encoding plasmid as template. A forward primer with a 5' portion overlapping with the artemin sequence and a reverse primer are then used to amplify the portion of the other, non-artemin growth factor member of the TGF-ß superfamily from the corresponding crossover point through the 3' end using a plasmid template containing the coding sequence for the non-artemin TGF-ß family member. The products of the two PCR reactions are gel purified and mixed together and a PCR reaction performed. Using an aliquot of this reaction as template a PCR reaction is performed using the artemin forward primer and the reverse primer for the non-artemin growth factor. The product is then cloned into an expression vector for production of the chimeric molecule.

Chimeric growth factors would be expected to be effective in promoting the growth and development of cells and for use in preventing the atrophy, degeneration or death of cells, particular in neurons. The chimeric polypeptides may also act as receptor antagonists of one or both of the full length growth factors from which the chimeric polypeptide was constructed or as an antagonist of any other growth factor that acts at the same receptor or receptors.

The present invention also includes therapeutic or pharmaceutical compositions comprising an artemin polypeptide in an effective amount for providing trophic support to cells in patients with cellular degeneration or dysfunction and a method comprising administering a therapeutically effective amount of the artemin polypeptide to a cell *ex vivo* or *in vivo*. The term "trophic support" is used herein to mean that a growth factor such as artemin provides sufficient nourishment to a cell such that the cell maintains or recovers at least one or more of its normal functions.

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The compositions and methods of the present invention are useful for treating a number of degenerative diseases and anaplastic diseases. Where the cellular degeneration, dysfunction or anaplasia involves neurons, the diseases include, but are not limited to peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease, Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, nervous system tumors such as neuroblastomas, multiple sclerosis, peripheral nerve trauma or injury, exposure to neurotoxins, metabolic diseases such as diabetes or renal dysfunctions and damage caused by infectious agents. In addition, artemin compositions can be used to treat enteric diseases such as idiopathic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers. If the cellular degeneration or dysfunction involves nonneuronal cells such as bone marrow cells, artemin may be useful in treating diseases including, but not limited to disorders of insufficient blood cells such as, for example, leukopenias including eosinopenia and/or basopenia, lymphopenia, monocytopenia, neutropenia, anemias, thrombocytopenia as well as an insufficiency of stem cells for any of the above. The cellular degeneration or dysfunction can also involve myocardial muscle cells in diseases such as cardiomyopathy and congestive heart failure. In addition small cell lung carcinoma can be treated with artemin polypeptide or polynucleotide compositions.

PCT/US99/22604

Treatment of enteric diseases with artemin includes the treatment of enteric neuropathies. The enteric nervous system is a complex collection of nerves that control the function of the gastrointestinal system, including gastrointestinal motility. Initial clinical studies with the NT-3, have shown that this neurotrophic factor increases gastrointestinal motility in normal volunteers and in patients suffering from peripheral neuropathies. Similarly, it is believed that artemin as well as the other members of the GDNF/neurturin/persephin family of growth factors will also show activity on enteric neurons. As a result, it is believed that artemin will be useful in treating enteric neuropathies such as in patients suffering from severe idiopathic constipation as well as patients suffering from constipation associated with Parkinson's disease, spinal cord injury, use of opiate pain-killers, and the like.

Whether artemin would be effectivity in the treatment of a particular cell type or tissues can be readily determined by one skilled in the art using any of a variety of

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assays known in the art. For example, with respect to providing trophic support for cells, trophic factors can produce beneficial biochemical and morphological effects and, under some circumstances, a promote cell survival. With respect to neurons, it is known in the art that depriving a neuron of trophic support results in a decrease in metabolic activity, i.e., glucose uptake, RNA synthesis and protein synthesis, required for normal function and growth. Deckwerth and Johnson, *J. Cell Biol. 123*:1207-1222, 1993. Removal of trophic support also results in a reduction in size of the cell body of the neuron. Presumably as a consequence of the loss of the metabolic effects of trophic factors, trophic factor deprivation results in a decrease or cessation of process outgrowth and may result in retraction of neuronal processes. In addition to the requirement of trophic factor for these aspects of neuronal biology, the neuron may require the neurotrophic factor to maintain survival; thus, survival assays are a frequently used means to detect or quantitate the actions of a neurotrophic factor. However, trophic support can also be manifest as morphological, biochemical, and functional changes; independent of neuronal number or any effect on survival.

As discussed above, growth factors can produce a cell differentiation in addition to providing trophic support for cells. Thus, it is believed that artemin polypeptides and polynucleotides can be beneficially used to produce a differentiation of anaplastic cells such as cancer cells. In particular, artemin can be used to treat nervous system tumors such as neuroblastomas. In addition, small cell lung carcinomas are known to express RET. Hence, it is believed that artemin can also be used to treat small cell lung carcinomas.

It is also contemplated that the eliciting of trophic support and/or differentiation can be achieved by administering an artemin polypeptide along with a GFRα3 polypeptide or by administering an artemin polynucleotide and a GFRα3 polynucleotide using the methods described herein for administration of artemin polypeptides or artemin polynucleotides. As noted above, Mouse GFRα3 was first identified as an expressed sequence tag (EST) that is homologous to GFRα1 and GFRα2 (Baloh et al., 1998, supra; Jing et al., 1997, supra; Naveilhan et al., Proc Natl Acad Sci USA 95, 1295-300, 1998; Widenfalk et al., Eur. J. Neurosci. 10, 1508-1517, 1998; Worby et al., J Biol Chem 273, 3502-3508, 1998). Human GFRα3 was identified by using the human GFRα2 amino acid sequence as a query to search a

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database of Expressed Sequence Tags (dbEST database) with the BLAST search algorithm (Altschul et al. J.Mol.Biol. 215:403-410, 1990; see copending application entitled "GFRa3, A Novel Member of the GDNF Coreceptor Family" which is file concurrently with the present application on December 22, 1998 and which is incorporated in its entirety herein by reference). Of the EST sequences identified, 5 several (AA049894, AA050083, AA041935, AA238748) did not correspond identically to either GFR $\alpha$ 1 or GFR $\alpha$ 2 but had significant homologies to both sequences. The clones corresponding to these EST's were acquired from the Washington University EST project and sequenced. One of these corresponded to a full-length mouse cDNA which the inventors herein named GFRa3. Sequence 10 information from the mouse cDNA was used to identify human genomic and cDNA clones, and the corresponding human and mouse predicted protein sequences are shown in Figure 12. The predicted amino acid sequences for human and mouse GFRα3 indicate a ~38.8 kDa protein with a putative N-terminal signal sequence 15 (amino acids 1-31 in hGFRα3 and 1-28 in mGFRα3 of Figure 12) (Nielsen et al., Protein Eng. 10:106, 1997), a mature GFRα3 polypeptide (amino acids 32-372 in hGFRα3 and 29-369 in mGFRα3 of Figure 12), three putative N-linked glycosylation sites, and a hydrophobic stretch of residues at the C-terminus consistent with a GPI signal peptide (amino acids 373-400 of hGFRα3 and 370-397 of mGFRα3 of Figure 12) (Undenfriend et al., Annu. Rev. Biochem. 64:563-591, 1995) like those 20 present in the related proteins GFRa1 and GFRa2 (Klein et al., Nature 387: 717-721, 1997; Jing et al., Cell 85:1113-1124, 1996; Treanor et al., Nature 382:80-83, 1996; Baloh et al., Neuron 18:793-802, 1997.). The human nucleotide sequence encoding GFRα3 precursor (SEQ ID NO:65) is shown in Figure 13. This precursor sequence includes the encoding sequence for the N-terminal signal sequence (nucleotides 1-93 25 of Figure 13), the encoding sequence for mature GFRa3 (nucleotides 94-1116 of Figure 13) and the encoding sequence for the C-terminal hydrophobic stretch of residues consistent with a GPI signal peptide (nucleotides 1117-1203 of Figure 13).

As described below, artemin binds to  $GFR\alpha 3$  in the absence of RET and it is believed that the resulting ligand/coreceptor complex is capable of binding to and activating RET receptors expressed by a target cell. Thus, treatment with artemin and

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GFR $\alpha$ 3 would be expected to increase the sensitivity of cells normally responsive to artemin and would also be expected to provide trophic support to cells that express RET but that are not normally responsive to artemin. Preferably, the artemin and GFR $\alpha$ 3 polypeptides are from the same species, i.e., human. It is also preferred that the GFR $\alpha$ 3 polypeptide be in soluble form, i.e., that it lack the GPI linkage to avoid potential undesirable interactions with cell membranes. As used herein a GFR $\alpha$ 3 polypeptide is intended to include the mature protein with or without the GPI anchor, as well as GFR $\alpha$ 3 fragments, particularly soluble fragments lacking a GPI anchor, that are capable of binding to both artemin and RET with such binding leading to the activation of RET.

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In certain circumstances, it may be desirable to modulate or decrease the amount of artemin expressed. Thus, in another aspect of the present invention, artemin anti-sense oligonucleotides can be made and a method utilized for diminishing the level of expression of artemin, respectively, by a cell comprising administering one or more artemin anti-sense oligonucleotides. By artemin anti-sense oligonucleotides reference is made to oligonucleotides that have a nucleotide sequence that interacts through base pairing with a specific complementary nucleic acid sequence involved in the expression of artemin such that the expression of artemin is reduced. Preferably, the specific nucleic acid sequence involved in the expression of artemin is a genomic DNA molecule or mRNA molecule that contains sequences of the artemin gene. Thus, the invention contemplates artemin anti-sense oligonucleotides that can base pair to flanking regions of the artemin gene, untranslated regions of artemin mRNA, the pre- or pro- portions of the artemin gene or the coding sequence for mature artemin protein. The term complementary to a nucleotide sequence in the context of artemin antisense oligonucleotides and methods therefor means sufficiently complementary to such a sequence as to allow hybridization to that sequence in a cell, i.e., under physiological conditions. The artemin antisense-oligonucleotides preferably comprise a sequence containing from about 8 to about 100 nucleotides and more preferably the artemin antisense oligonucleotides comprise from about 15 to about 30 nucleotides. The artemin antisense oligonucleotides can also contain a variety of modifications that confer resistance to nucleolytic degradation such as, for example, modified internucleoside

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linkages (Uhlmann and Peyman, *Chemical Reviews 90*:543-548, 1990; Schneider and Banner, *Tetrahedron Lett 31*:335, 1990), modified nucleic acid bases and/or sugars and the like.

The therapeutic or pharmaceutical compositions of the present invention can be administered by any suitable route known in the art including for example intravenous, subcutaneous, intramuscular, transdermal, intrathecal or intracerebral. Administration can be either rapid as by injection or over a period of time as by slow infusion or administration of slow release formulation. For treating tissues in the central nervous system, administration can be by injection or infusion into the cerebrospinal fluid (CSF). When it is intended that artemin be administered to cells in the central nervous system, administration can be with one or more agents capable of promoting penetration of artemin across the blood-brain barrier.

Artemin can also be linked or conjugated with agents that provide desirable pharmaceutical or pharmacodynamic properties. For example, artemin can be coupled to any substance known in the art to promote penetration or transport across the blood-brain barrier such as an antibody to the transferrin receptor, and administered by intravenous injection. (See for example, Friden et al., *Science 259*:373-377, 1993). Furthermore, artemin can be stably linked to a polymer such as polyethylene glycol to obtain desirable properties of solubility, stability, half-life and other pharmaceutically advantageous properties. (See for example Davis et al. *Enzyme Eng 4:*169-73, 1978; Burnham, *Am J Hosp Pharm 51*:210-218, 1994). Preferably, artemin is administered with a carrier such as liposomes or polymers containing a targeting moiety to limit delivery of artemin to targeted cells. Examples of targeting moieties include but are not limited to antibodies, ligands or receptors to specific cell surface molecules.

For nonparenteral administration, the compositions can also include absorption enhancers which increase the pore size of the mucosal membrane. Such absorption enhancers include sodium deoxycholate, sodium glycocholate, dimethyl-β-cyclodextrin, lauroyl-1-lysophosphatidylcholine and other substances having structural similarities to the phospholipid domains of the mucosal membrane.

The compositions are usually employed in the form of pharmaceutical preparations. Such preparations are made in a manner well known in the

pharmaceutical art. One preferred preparation utilizes a vehicle of physiological saline solution, but it is contemplated that other pharmaceutically acceptable carriers such as physiological concentrations of other non-toxic salts, five percent aqueous glucose solution, sterile water or the like may also be used. It may also be desirable that a suitable buffer be present in the composition. Such solutions can, if desired, be lyophilized and stored in a sterile ampoule ready for reconstitution by the addition of sterile water for ready injection. The primary solvent can be aqueous or alternatively non-aqueous. Artemin can also be incorporated into a solid or semi-solid biologically compatible matrix which can be implanted into tissues requiring treatment.

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The carrier can also contain other pharmaceutically-acceptable excipients for modifying or maintaining the pH, osmolarity, viscosity, clarity, color, sterility, stability, rate of dissolution, or odor of the formulation. Similarly, the carrier may contain still other pharmaceutically-acceptable excipients for modifying or maintaining release or absorption or penetration across the blood-brain barrier. Such excipients are those substances usually and customarily employed to formulate dosages for parenteral administration in either unit dosage or multi-dose form or for direct infusion into the cerebrospinal fluid by continuous or periodic infusion.

Dose administration can be repeated depending upon the pharmacokinetic parameters of the dosage formulation and the route of administration used.

It is also contemplated that certain formulations containing artemin are to be administered orally. Such formulations are preferably encapsulated and formulated with suitable carriers in solid dosage forms. Some examples of suitable carriers, excipients, and diluents include lactose, dextrose, sucrose, sorbitol, mannitol, starches, gum acacia, calcium phosphate, alginates, calcium silicate, microcrystalline cellulose, polyvinylpyrrolidone, cellulose, gelatin, syrup, methyl cellulose, methyl- and propylhydroxybenzoates, talc, magnesium, stearate, water, mineral oil, and the like. The formulations can additionally include lubricating agents, wetting agents, emulsifying and suspending agents, preserving agents, sweetening agents or flavoring agents. The compositions may be formulated so as to provide rapid, sustained, or delayed release of the active ingredients after administration to the patient by employing procedures well known in the art. The formulations can also contain

substances that diminish proteolytic degradation and promote absorption such as, for example, surface active agents.

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The specific dose is calculated according to the approximate body weight or body surface area of the patient or the volume of body space to be occupied. The dose will also be calculated dependent upon the particular route of administration selected. Further refinement of the calculations necessary to determine the appropriate dosage for treatment is routinely made by those of ordinary skill in the art. Such calculations can be made without undue experimentation by one skilled in the art based on the activity of artemin for a particular cell type *in vitro*. The activity of artemin on various peripheral and central neurons in culture is described below and artemin activity on a particular target cell type can be determined by routine experimentation. Exact dosages are determined in conjunction with standard doseresponse studies. It will be understood that the amount of the composition actually administered will be determined by a practitioner, in the light of the relevant circumstances including the condition or conditions to be treated, the choice of composition to be administered, the age, weight, and response of the individual patient, the severity of the patient's symptoms, and the chosen route of administration.

In one embodiment of this invention, artemin may be therapeutically administered by implanting into patients vectors or cells capable of producing a biologically-active form of artemin or a precursor of artemin, i.e. a molecule that can be readily converted to a biological-active form of artemin by the body. In one approach cells that secrete artemin may be encapsulated into semipermeable membranes for implantation into a patient. The cells can be cells that normally express artemin or a precursor of artemin or the cells can be transformed to express artemin or a precursor thereof. In some embodiments, the cells are transformed to express and secrete both artemin and GFR $\alpha$ 3, preferably in a soluble form. It is preferred that the cell be of human origin and that the artemin be human artemin when the patient is human. However, the formulations and methods herein can be used for veterinary as well as human applications and the term "patient" as used herein is intended to include human and veterinary patients.

Cells can be grown ex vivo for use in transplantation or engraftment into patients (Muench et al., Leuk & Lymph 16:1-11, 1994). In another embodiment of the

present invention, artemin can be used to promote the ex vivo expansion of a cells for transplantation or engraftment. Current methods have used bioreactor culture systems containing factors such as erythropoietin, colony stimulating factors, stem cell factor, and interleukins to expand hematopoietic progenitor cells for erythrocytes, monocytes, neutrophils, and lymphocytes (Verfaillie, Stem Cells 12:466-476, 1994). 5 These stem cells can be isolated from the marrow of human donors, from human peripheral blood, or from umbilical cord blood cells. The expanded blood cells are used to treat patients who lack these cells as a result of specific disease conditions or as a result of high dose chemotherapy for treatment of malignancy (George, Stem Cells 12(Suppl 1):249-255, 1994). In the case of cell transplant after chemotherapy, 10 autologous transplants can be performed by removing bone marrow cells before chemotherapy, expanding the cells ex vivo using methods that also function to purge malignant cells, and transplanting the expanded cells back into the patient following chemotherapy (for review see Rummel and Van Zant, J Hematotherapy 3:213-218, 15 1994).

It is also believed that artemin with or without GFR $\alpha$ 3 can be used for the *ex vivo* expansion of precursor cells in the nervous system. Transplant or engraftment of cells is currently being explored as a therapy for diseases in which certain populations of neurons are lost due to degeneration such as, for example, in Parkinson's disease (Bjorklund, *Curr Opin Neurobiol 2*:683-689, 1992). Neuronal precursor cells can be obtained from animal or human donors or from human fetal tissue and then expanded in culture using artemin. These cells can then be engrafted into patients where they would function to replace some of the cells lost due to degeneration. Because neurotrophins have been shown to be capable of stimulating the survival and proliferation of neuronal precursor cells such as, for example, NT-3 stimulation of sympathetic neuroblast cells (Birren et al., *Develop 119*:597-610, 1993), artemin could also function in similar ways during the development of the nervous system and could be useful in the *ex vivo* expansion of neuronal cells.

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In a number of circumstances it would be desirable to determine the levels of artemin or soluble GFR $\alpha$ 3 in a patient. A change in the amount of endogenously produced artemin or GFR $\alpha$ 3 may play a role in certain disease conditions, particularly where there is cellular degeneration such as in neurodegenerative conditions or

diseases. Other neurotrophic factors are known to change during disease conditions. For example, in multiple sclerosis, levels of NGF protein in the cerebrospinal fluid are increased during acute phases of the disease (Bracci-Laudiero et al., *Neuroscience Lett 147*:9-12, 1992) and in systemic lupus erythematosus there is a correlation between inflammatory episodes and NGF levels in sera (Bracci-Laudiero et al. *NeuroReport 4*:563-565, 1993).

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Thus, quantification of artemin levels may provide clinically useful information. Furthermore, in the treatment of degenerative conditions, compositions containing artemin can be administered and it would likely be desirable to achieve certain target levels of artemin in sera, in cerebrospinal fluid or in any desired tissue compartment. It would, therefore, be advantageous to be able to monitor the levels of artemin in a patient. Accordingly, the present invention also provides methods for detecting the presence of artemin in a sample from a patient.

The term "detection" as used herein in the context of detecting the presence of artemin in a patient is intended to include the determining of the amount of artemin or the ability to express an amount of artemin in a patient, the distinguishing of artemin from other growth factors, the estimation of prognosis in terms of probable outcome of a degenerative disease and prospect for recovery, the monitoring of the artemin levels over a period of time as a measure of status of the condition, and the monitoring of artemin levels for determining a preferred therapeutic regimen for the patient.

To detect the presence of artemin in a patient, a sample is obtained from the patient. The sample can be a tissue biopsy sample or a sample of blood, plasma, serum, CSF or the like. Samples for detecting artemin can be taken from any tissue known to express artemin. When assessing peripheral levels of artemin, it is preferred that the sample be a sample of blood, plasma or serum or alternatively from a tissue biopsy sample. When assessing the levels of artemin in the central nervous system a preferred sample is a sample obtained from cerebrospinal fluid.

In some instances it is desirable to determine whether the artemin gene is intact in the patient or in a tissue or cell line within the patient. By an intact artemin gene it is meant that there are no alterations in the gene such as point mutations, deletions, insertions, chromosomal breakage, chromosomal rearrangements and the

like wherein such alteration might alter production of artemin or alter its biological activity, stability or the like to lead to disease processes or susceptibility to cellular degenerative conditions. Conversely, by a non-intact artemin gene it is meant that such alterations are present. Thus, in one embodiment of the present invention a method is provided for detecting and characterizing any alterations in the artemin gene. The method comprises providing a polynucleotide that specifically hybridizes to an artemin cDNA, genomic DNA or a fragment thereof.

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Typically, patient genomic DNA is isolated from a cell sample from the patient and digested with one or more restriction endonucleases such as, for example, TaqI and AluI. Using the Southern blot protocol, which is well known in the art, this assay determines whether a patient or a particular tissue in a patient has an intact artemin gene or an abnormality in the artemin gene. Hybridization to the artemin gene would involve denaturing the chromosomal DNA to obtain a single-stranded DNA; contacting the single-stranded DNA with a gene probe associated with the artemin gene sequence; and identifying the hybridized DNA-probe to detect chromosomal DNA containing at least a portion of the human artemin gene.

The term "probe" as used herein refers to a structure comprised of a polynucleotide which forms a hybrid structure with a target sequence, due to complementarity of probe sequence with a sequence in the target region. The probes need not contain the exact complement of the target sequence, but must be sufficiently complementary to selectively hybridize with the strand being detected. By selective hybridization or specific hybridization it is meant that a polynucleotide preferentially hybridizes to a target polynucleotide. Oligomers suitable for use as probes may contain a minimum of about 8-12 contiguous nucleotides which are complementary to the targeted sequence and preferably a minimum of about 15 or 17 nucleotides although polynucleotide probes of about 20 to 25 nucleotides and up to about 100 nucleotides or even greater are within the scope of this invention.

The artemin gene probes of the present invention can be DNA or RNA oligonucleotides and can be made by any method known in the art such as, for example, excision, transcription or chemical synthesis. Probes may be labeled with any detectable label known in the art such as, for example, radioactive or fluorescent labels or enzymatic marker. Labeling of the probe can be accomplished by any

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method known in the art such as by PCR, random priming, end labeling, nick translation or the like. One skilled in the art will also recognize that other methods not employing a labeled probe can be used to determine the hybridization. Examples of methods that can be used for detecting hybridization include Southern blotting, fluorescence *in situ* hybridization, and single-strand conformation polymorphism with PCR amplification.

Hybridization is typically carried out at 25-45°C, more preferably at 32-40°C and more preferably at 37-38°C. The time required for hybridization is from about 0.25 to about 96 hours, more preferably from about one to about 72 hours, and most preferably from about 4 to about 24 hours.

Artemin gene abnormalities can also be detected by using the PCR method or any other known DNA amplification method, which uses oligonucleotides to identify a target sequence within a longer sequence, and primers that flank or lie within the artemin gene. The PCR method is well known in the art. Briefly, this method is performed using two oligonucleotide primers which are capable of hybridizing to the nucleic acid sequences flanking a target sequence that lies within an artemin gene and amplifying the target sequence. The terms "oligonucleotide primer" as used herein refers to a short strand of DNA or RNA typically ranging in length from about 8 to about 30 bases. The upstream and downstream primers are preferably a minimum of from about 15 nucleotides to about 20 nucleotides and up to about 30 nucleotides or even greater in length. The primers can hybridize to the flanking regions for replication of the target nucleotide sequence. The polymerization is catalyzed by a DNA-polymerase in the presence of deoxynucleotide triphosphates or nucleotide analogs to produce double-stranded DNA molecules. The double strands are then separated by any denaturing method including physical, chemical or enzymatic. Commonly, the method of physical denaturation is used involving heating the nucleic acid, typically to temperatures from about 80°C to 105°C for times ranging from about 1 to about 10 minutes. The process is repeated for the desired number of cycles.

The primers are selected to be substantially complementary to the strand of DNA being amplified. Therefore, the primers need not reflect the exact sequence of the template, but must be sufficiently complementary to selectively hybridize or

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specifically hybridize with the strand being amplified. By selective hybridization or specific hybridization it is meant that a polynucleotide preferentially hybridizes to a target polynucleotide.

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After PCR amplification, the DNA sequence comprising an artemin-encoding nucleotide sequence or a fragment thereof is then directly sequenced and analyzed by comparison of the sequence with the sequences disclosed herein to identify alterations which might change activity or expression levels or the like.

In another embodiment a method for detecting artemin is provided based upon an analysis of tissue expressing the artemin gene. The method comprises hybridizing a polynucleotide probe to mRNA from a sample of tissues that normally express the artemin gene or from a cDNA produced from the mRNA of the sample. The sample is obtained from a patient suspected of having an abnormality in the artemin gene or from a particular patient tissue or cell type suspected of having an abnormality in the artemin gene.

To detect the presence of mRNA encoding artemin protein, a sample is obtained from a patient. The sample can be from blood or from a tissue biopsy sample. The sample may be treated to extract the mRNA contained therein. The resulting mRNA from the sample is subjected to gel electrophoresis or other size separation techniques.

The mRNA of the sample is contacted with a nucleic acid serving as a probe to form hybrid duplexes. The use of a labeled probes as discussed above allows detection of the resulting duplex.

High stringency conditions can be used in order to prevent false positives, that is hybridization to non-artemin nucleotide sequences. When using sequences that are not perfectly complementary to an artemin-encoding polynucleotide or a fragment thereof, less stringent conditions could be used, however, this would be a less preferred approach because of the likelihood of false positives. The stringency of hybridization is determined by a number of factors during hybridization and during the washing procedure, including temperature, ionic strength, length of time and concentration of formamide. These factors are outlined in, for example, Sambrook et al. (Sambrook, et al., 1989, *supra*).

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In order to increase the sensitivity of the detection in a sample of mRNA encoding the artemin protein, the technique of reverse transcription/polymerization chain reaction (RT/PCR) can be used to amplify cDNA transcribed from mRNA encoding the artemin protein. The method of RT/PCR is well known in the art.

Alternatively, an artemin target sequence in the reverse transcribed cDNA can be amplified and detected using any other known methodology such as ligase chain reaction methods, including gap LCR (G-LCR) and other variations, or self-sustained sequence replication (3SR) and its various modifications. In addition, the artemin mRNA can be detected directly by asymmetric gap LCR (AG-LCR). See, e.g., Leckie et al., "Infectious Disease Testing by Ligase Chain Reaction" in *Molecular Biology and Biotechnology*, R. A. Myers, ed., pp. 463-466, VCH Publishers, 1995.

The present invention further provides for methods to detect the presence of the artemin protein in a sample obtained from a patient. Any method known in the art for detecting proteins can be used. Such methods include, but are not limited to immunodiffusion, immunoelectrophoresis, immunochemical methods, binder-ligand assays, immunohistochemical techniques, agglutination and complement assays. (for example see *Basic and Clinical Immunology*, Sites and Terr, eds., Appleton & Lange, Norwalk, Conn. pp 217-262, 1991). Preferred are binder-ligand immunoassay methods including reacting antibodies with an epitope or epitopes of the artemin protein and competitively displacing the artemin protein.

Numerous competitive and non-competitive protein binding immunoassays are well known in the art. Antibodies employed in such assays may be unlabeled, for example as used in agglutination tests, or labeled for use in a wide variety of assay methods. Labels that can be used include radionuclides, enzymes, fluorescers, chemiluminescers, enzyme substrates or co-factors, enzyme inhibitors, particles, dyes and the like for use in radioimmunoassay (RIA), enzyme immunoassays, e.g., enzyme-linked immunosorbent assay (ELISA), fluorescent immunoassays and the like.

Polyclonal or monoclonal antibodies to the artemin protein or to an epitope thereof can be made for use in immunoassays by any of a number of methods known in the art. By epitope reference is made to an antigenic determinant of a polypeptide. The term epitope can also include artemin-specific B cell epitopes or T helper cell

epitopes. An epitope could comprise 3 amino acids in a spatial conformation which is unique to the epitope. Generally an epitope consists of at least 6 such amino acids. Methods of determining the spatial conformation of amino acids are known in the art, and include, for example, x-ray crystallography and 2 dimensional nuclear magnetic resonance.

One approach for preparing antibodies to a protein is the selection and preparation of an amino acid sequence of all or part of the protein, chemically synthesizing the sequence and injecting it into an appropriate animal, usually a rabbit or a mouse (See Example 10).

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Oligopeptides can be selected as candidates for the production of an antibody to the artemin protein based upon the oligopeptides lying in hydrophilic regions, which are thus likely to be exposed in the mature protein.

Antibodies to artemin can also be raised against oligopeptides that include one or more of the conserved regions identified herein such that the antibody can cross-react with other family members. Such antibodies can be used to identify and isolate the other family members.

Methods for preparation of the artemin protein or an epitope thereof include, but are not limited to chemical synthesis, recombinant DNA techniques or isolation from biological samples. Chemical synthesis of a peptide can be performed, for example, by the classical Merrifeld method of solid phase peptide synthesis (Merrifeld, *J Am Chem Soc 85:*2149, 1963) or the FMOC strategy on a Rapid Automated Multiple Peptide Synthesis system (DuPont Company, Wilmington, DE) (Caprino and Han, *J Org Chem 37:*3404, 1972).

Polyclonal antibodies can be prepared by immunizing rabbits or other animals by injecting antigen followed by subsequent boosts at appropriate intervals. The animals are bled and sera assayed against purified artemin protein usually by ELISA or by bioassay based upon the ability to block the action of artemin. When using avian species, e.g. chicken, turkey and the like, the antibody can be isolated from the yolk of the egg. Monoclonal antibodies can be prepared after the method of Milstein and Kohler by fusing splenocytes from immunized mice with continuously replicating tumor cells such as myeloma or lymphoma cells. (Milstein and Kohler *Nature* 256:495-497, 1975; Gulfre and Milstein, *Methods in Enzymology: Immunochemical* 

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Techniques 73:1-46, Langone and Banatis eds., Academic Press, 1981). The hybridoma cells so formed are then cloned by limiting dilution methods and supernates assayed for antibody production by ELISA, RIA or bioassay.

The unique ability of antibodies to recognize and specifically bind to target proteins provides an approach for treating an over expression of the protein. Thus, another aspect of the present invention provides for a method for preventing or treating diseases involving over expression of the artemin protein by treatment of a patient with specific antibodies to the artemin protein.

Specific antibodies, either polyclonal or monoclonal, to the artemin protein can be produced by any suitable method known in the art as discussed above. For example, murine or human monoclonal antibodies can be produced by hybridoma technology or, alternatively, the artemin protein, or an immunologically active fragment thereof, or an anti-idiotypic antibody, or fragment thereof can be administered to an animal to elicit the production of antibodies capable of recognizing and binding to the artemin protein. Such antibodies can be from any class of antibodies including, but not limited to IgG, IgA, IgM, IgD, and IgE or in the case of avian species, IgY and from any subclass of antibodies.

Preferred embodiments of the invention are described in the following examples. Other embodiments within the scope of the claims herein will be apparent to one skilled in the art from consideration of the specification or practice of the invention as disclosed herein. It is intended that the specification, together with the examples, be considered exemplary only, with the scope and spirit of the invention being indicated by the claims which follow the examples.

25 Example 1

This example illustrates the identification and cloning of polynucleotides encoding human and mouse artemin.

Artemin was discovered by using the full length murine pre-pro neurturin amino acid sequence (195 amino acids) to scan the High Throughout Genome Sequences (htgs) database. The database was searched using the tblastn feature of the BLAST 2.0.5 program using the default parameters (Altschul, et al., Nucleic Acids Res. 25:3389-3402, 1997). This search identified two human bacterial artificial

chromosome (BAC) clones, AC005038 (NH0486I22) and AC005051 (RG037F03), with significant homology scores when aligned with murine pre-pro neurturin. When the BLAST search was repeated using the mature human neurturin sequence as a query, the same two BAC clones were identified. Using the sequences of these two BAC clones, two EST's (Expressed Sequence Tags) were also identified in the EST database using the blastn feature of BLAST 2.0.5 program using the default parameters. These ESTs were: AA931637 (0035c12) from a lung carcinoma and AA533512 (nj96b11) from a prostate carcinoma. Analysis of these alignments indicated that portions of the anonymous htgs sequences possibly represented a new member of the GDNF-neurturin-persephin family. Indeed, although, the hgts sequences were later discovered to have numerous sequence errors, i.e. omissions, additions and incorrect bases, one of the BAC clones, AC005038, had a stretch of 197 nucleotides (nucleotides 67,464 to 67,660) which ultimately turned out to be identical to nucleotides 663-467 of the complementary strand of the artemin nucleic acid (Fig. 1B) and the other, AC005051, had a stretch of 183 nucleotides (nucleotides 113,379 to 113,561) identical to nucleotides 648-468 of the complementary strand of the artemin nucleic acid (Fig. 1B).

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Because sequences in the hgts database were generally known to contain many sequence errors, oligonucleotide primers were designed from the sequences of the two 20 htgs entries to obtain fragments of the gene for this new growth factor from human genomic DNA. The primers were designed to amplify an approximate 708 nt fragment encoding a portion of the predicted new growth factor, which would extend from within the pro-domain, through the mature region of the protein, and end within the 3' untranslated region. Human genomic DNA was used as a template in a PCR 25 reaction which utilized a forward primer M4206 [5'-TGGCCGCTCTGGCTCTGAGCA-3'] (SEQ ID NO:20) and reverse primer M4205 [5'-CGATCATCTAGACCACCGGTAAGGGTCCAGTCTGCAA-3'] (SEQ ID NO:21). The PCR reaction was carried out using Klentag in Klentag buffer with additives (1.5 M betaine/1.5% DMSO; final concentration) and the following parameters: initial denaturation at 95°C for 5 min, then 95°C for 15 sec, 60°C for 30 30 sec, 68°C for 80 sec x 30 cycles. The amplified products were kinased with T4 polynucleotide kinase, the ends were blunted with E. coli DNA polymerase I (Klenow

fragment), and cloned into EcoRV digested BSKS plasmid. The nucleotide sequences of the resulting clones were obtained.

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An additional, overlapping fragment was also amplified from human genomic DNA, which extends from the beginning of the mature region to the stop codon. This was amplified by PCR using the forward primer M4207 (5'-TACAAGGACGATGACGATAAGGGGGCGCGGGGCTGCGCCTCGCCAGCTCGCCAGCTGCGCTCGCCAGCTGCGCTCGCCAGCTGCGCTCGCCAGCTGCGCTCGCCAGTCTGCAA-3') (SEQ ID NO:23). The PCR reaction was carried out using Klentaq in Klentaq buffer with additives (5% DMSO: final concentration) and the following parameters: initial denaturation at 95°C

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DMSO; final concentration) and the following parameters: initial denaturation at 95°C for 5 min, then 95°C for 15 sec, 60°C for 30 sec, 68°C for 80 sec x 30 cycles. The amplified products were kinased with T4 polynucleotide kinase, the ends were blunted with E. coli DNA polymerase I (Klenow fragment), and cloned into EcoRV digested BSKS plasmid. The nucleotide sequences of the resulting clones were obtained.

The sequences from these fragments and the sequences of the entries in the htgs database were assembled using Seqman program (DNASTAR). The assembled nucleotide sequence, which is shown in Fig. 1A, encodes a partial pro-artemin amino acid sequence which extends from within the pro-domain, includes 3 potential RXXR cleavage sites and then includes a region corresponding to the mature artemin polypeptide, which is homologous to mature GDNF, neurturin and persephin as shown in Fig. 2A.

Primers designed from this sequence were also used to PCR amplify a fragment of mouse genomic DNA which corresponded to the mouse artemin gene. This mouse DNA fragment was than used to screen mouse BAC genomic libraries and a BAC clone containing the mouse artemin gene was identified (data not shown).

Primers were made from the human and mouse artemin genomic sequences to perform rapid amplification of cDNA ends (RACE) PCR to obtain the 5' end of the artemin cDNA from several tissue libraries. PCR fragments were blunt-cloned into EcoRV digested BSKS as above, and sequenced entirely. Human RACE products were obtained from MARATHON RACE cDNA libraries (CLONTECH) prepared from pituitary, placenta, and kidney. Mouse RACE products were obtained from an

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embryonic day 18 (E18) mouse MARATHON RACE library. Complete double-stranded sequence analysis of cloned PCR fragments from genomic DNA and cDNA was performed to generate contigs of the human and mouse cDNAs. At least two independent clones of each PCR fragment were sequenced. Genomic sequence of mouse BAC clone restriction fragments (subcloned into pBluescript), and sequences derived from portions of the human BAC clone, were used to confirm the cDNA sequences, which are shown in Fig. 1B (human) and Fig. 1C (mouse).

Intron locations in the cDNA and splicing were confirmed by sequencing PCR amplified fragments from human and mouse cDNA libraries that cross both introns. As illustrated in Fig. 2C, the artemin gene has two introns. cDNAs containing a second starting methionine (Met\*) produced by alternative splicing of the first intron were also identified by RACE PCR from some human cDNA libraries; however, a similar message was not identified in mouse, and the mouse genomic sequence does not contain a methionine in this position. Mature artemin protein generated from either of the human messages would be identical.

Analysis of markers present on the artemin-encoding BAC clones indicated that the human artemin gene is located on chromosome 1p32-33, flanked by markers D1S190-artemin-D1S211 (telomeric to centromeric).

The human DNA sequence as shown in Figure 14, which includes the Artemin gene, was compiled from sequences from multiple independently cloned PCT products using human genomic DNA as template and from the human BAC clone sequences. The beginning of the sequence corresponds to nt 11472 of BAC clone AC005051, and nt 68862 of BAC clone AC005038 from the htgs database as described above.

A rat artemin cDNA fragment as shown in Figure 15 was generated using PCT from a rat Schwann cell cDNA library as template with the following primers: forward 5'-CCGGTGAGCGCTCTCGGCCT-3' (SEQ ID NO:76) and reverse primer 5'-TTCTGGATTCTCCCAGAGGAGTTC-3' (SEQ ID NO:77). PCT conditions were as follows: 95°C for 2 min; then 30 cycles of 95°C for 10 sec, 60°C for 30 sec; 72°C for 30 sec, then 72°C for 5 min;

WO 00/18799 PCT/US99/22604

# 52 **Example 2**

This example illustrates the expression of artemin in various human tissues.

As an initial survey of artemin expression in adult and embryonic human tissues, a human master RNA blot (MRB) (CLONTECH) containing normalized samples of poly(A)RNA was probed with a random hexamer <sup>32</sup>P-labeled fragment of the human artemin cDNA and signals were visualized using a PhosphorImager and (Molecular Dynamics). The results are shown in Fig. 5A.

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Relatively low level expression was observed in many adult tissues, with the highest level in the pituitary gland, placenta and trachea. Among human fetal tissues, kidney and lung showed the highest level of expression. While little signal was visible in the adult and fetal brain, low-level expression of artemin mRNA was observed in structures of the adult basal ganglia (subthalamic nucleus, putamen, substantia nigra) and in the adult thalamus, suggesting artemin may influence subcortical motor systems. Expression was also observed in the spinal cord, however the RNA preparation from this tissue contains DRG material as well as spinal cord, which likely contributes to the observed expression (see Figure 6).

To more precisely localize artemin expression during embryogenesis, a fragment of the rat artemin cDNA was generated using PCR and cloned into pBluescript. Sense and anti-sense <sup>33</sup>P-labeled RNA probes were generated from the cloned rat artemin cDNA fragment and used for *in situ* hybridization analysis of artemin mRNA expression in fresh-frozen embryonic day 14 (E14) rat embryos, which was performed as described previously (Araki et al., *Neuron 17*:353-361, 1998). The results are shown in Figs. 6A and 6B.

No signal was observed in the brain or spinal cord at this embryonic age. The most striking expression was observed in the nerve roots, but not the developing neurons, of the dorsal root ganglia (DRG) (Fig. 6A). Furthermore, significant diffuse expression was also observed surrounding the superior mesenteric artery, corresponding to either the surrounding mesenchyme or expression by cell of the artery itself (Fig. 6B). These data suggest the possibility that artemin may act as a survival/trophic factor for peripheral neurons, either in a paracrine fashion for developing sensory neurons of the DRG, or as a target-derived factor for autonomic innervation of the superior mesenteric artery. Interestingly, among potential GFRα

receptors, expression of artemin is most complementary with that of the orphan GFRα3, which is expressed at high levels in peripheral ganglia, but which has no detectable expression in the developing CNS by *in situ* hybridization (Baloh et al., 1998, *supra*; Trupp et al., *Mol. Cell. Neurosci.*, 11:47-63, 1998; Widenfalk et al., *Eur. J. Neurosci.* 10:1508-1517, 1998; Worby et al., *J. Biol. Chem.* 271, 23619-23622, 1998). No signal was detected using the sense probe on any tissue samples.

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The expression of artemin in the developing nerve roots at E14 suggested it is produced by Schwann cell precursors or immature Schwann cells. To investigate if Schwann cells express artemin, semi-quantitative RT-PCR analysis was performed on cDNA libraries prepared from primary cultures of Schwann cells isolated from early post-natal rats, from myelinating Schwann cells from the adult sciatic nerve and from Schwann cells isolated from the distal segment of the sciatic nerve at 16 hours, 3 days or 7 days following nerve transection. Cultures of purified Schwann cells from early post-natal rats was performed as described by Brockes et al., Brain Res. 165:105-118, 1979. Sciatic nerve transection in adult rats and generation of reverse-transcribed cDNA were performed as described elsewhere (Araki et al., supra; Baloh et al., Neuron 18:793-802, 1997). cDNA samples from the libraries were normalized to levels of glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and then amplified by RT-PCR using as forward primer 5'-TCGCGACGGTGGCTCACCGGTCTT-3' (SEO ID NO:66) and as reverse primer 5'-GCACGAGCCGCTGCAGAAGCGGAA-3' (SEQ ID NO:67) and the following conditions: 95°C for 2 min followed by 36 cycles of 95°C, 20 sec; 54°C, 30 sec; and 72°C, 30 sec.; and then a final incubation at 72°C, 2 min. Products were separated on a 3% agarose gel and stained with Ethidium Bromide.

As shown in Fig. 6C, artemin is expressed at much higher levels in immature Schwann cells in culture than by mature myelinating Schwann cells (PS) of the adult sciatic nerve (N). However, artemin expression is up-regulated in the distal nerve segment after sciatic transection, a paradigm in which Schwann cells reacquire an immature state to support regenerating axons (reviewed in Scherer, *Curr. Opin Neurol.* 10:386-397, 1997). These data indicate that Schwann cells produce artemin, and furthermore suggest that artemin expression is regulated appropriately to influence developing and regenerating peripheral neurons.

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#### Example 3

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This example illustrates the production of recombinant artemin protein.

PCT/US99/22604

A PCR fragment was generated which corresponded to the mature human artemin coding sequence (Fig. 3A), with an NdeI site and an 8-histidine tag on the 5' end, and a KpnI site on the 3' end, and then cloned directly into the corresponding sites of the pET30a(+) bacterial expression vector (Novagen). The artemin-encoding expression vector was transformed into the *E. coli* strain BL21 and recombinant human artemin protein was produced and purified as described previously for neurturin (Creedon et al., *Proc. Natl. Acad. Sci. USA* 94:7018-7023, 1997).

#### Example 4

This example illustrates that artemin promotes the survival of peripheral neurons *in vitro*.

As shown above, expression of artemin suggests it may influence developing peripheral neurons, like GDNF and neurturin. To assess artemin's ability to support the survival of different peripheral neuronal populations *in vitro*, neurons from the dorsal root, trigeminal, nodose, and superior cervical ganglia (SCG) derived from postnatal day 1 (P1) Sprague-Dawley rats (Harlan Sprague-Dawley, IN) were cultured in the presence of artemin, GDNF, neurturin or persephin. Each GDNF ligand family member consisted of the human mature amino acid sequence and was produced by standard recombinant DNA methods. Recombinant GDNF, neurturin and persephin were obtained from Genentech while recombinant artemin was produced as described in Example 3.

The cultures were performed using standard methods described elsewhere (Kotzbauer et al., *Neuron* 12:763-773, 1994; Kotzbauer et al., *Nature* 384:467-470, 1996; Milbrandt et al., *Neuron* 20:245-253, 1998). In brief, for SCG neurons, after dissection and dissociation, neurons were plated on collagen-coated 24-well tissue culture plates and maintained in NGF-containing medium (AM50) for 5 days, after which they were either kept in AM50 or switched to medium containing neutralizing anti-NGF antibodies plus GDNF, neurturin, persephin, artemin, (50 ng/ml where not noted otherwise), or no growth factors. Cultures were maintained for 3 days, after

which they were fixed in 4% paraformaldehyde and stained with toluidine blue, and surviving neurons were counted. For nodose ganglion and dorsal root ganglion neuron cultures, dissociated neurons were plated directly into NGF (for dorsal root ganglia), or BDNF (for nodose ganglia, 100ng/ml) or one of the GDNF ligands in serum-containing medium (AM50 with anti-NGF antibodies), and cells surviving after 3 days in culture were counted as above. Trigeminal ganglia were dissected, dissociated, and plated directly into medium containing NGF or the indicated GDNF ligands, and the number of surviving cells was assessed after 3 days *in vitro*. For all culture systems, 2-3 independent experiments were performed. The results are shown in Figs. 7A-7E.

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Like GDNF and neurturin (NTN), artemin (ART) supported the survival of a subset of sensory neurons from both the dorsal root ganglion (DRG) and the trigeminal ganglion (TG), whereas persephin (PSP) did not (Figure 7A and 7C). Interestingly, in both these populations of sensory neurons ART supported a larger number of neurons than GDNF or neurturin, and supported a similar (DRG) or a greater number (TG) of neurons than NGF. Dose response analysis of artemin's survival promoting effect on DRG neurons (Fig. 7B) revealed an EC<sub>50</sub> of 1-3 ng/ml, ~10<sup>-10</sup>M.

With respect to visceral sensory neurons of the nodose ganglion (NG), which consists of a large number of neurons responsive to the neurotrophin BDNF (Lindsay et al., 1985), artemin, GDNF, and neurturin supported similar numbers of NG neurons, and each factor yielded equal or greater survival promotion than BDNF (Fig. 7D). These results were similar to previous reports (Kotzbauer et al., 1996, *supra*; Trupp et al., 1995, *supra*).

Artemin also supported the survival of SCG neurons in culture (Fig. 7E), although fewer neurons were supported by artemin than by GDNF or neurturin. In contrast to the sensory neurons examined above, none of the GDNF family ligands supported as many neurons as NGF. Therefore, similar to GDNF and neurturin, and as consistent with its embryonic expression pattern, artemin is a survival factor for sensory and sympathetic peripheral neurons in culture.

WO 00/18799 PCT/US99/22604

# 56 Example 5

This example illustrates that RET-expressing neuroblastoma cell lines are responsive to artemin.

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Neuroblastoma cell lines are derivatives of peripheral sympathetic neuroblast tumors that respond to GDNF and neurturin and display a variety of GFRα/RET receptor profiles (Hishiki et al., *Cancer Res.* 58:2158-2165, 1998; Tansey et al., submitted). Artemin's ability to influence several of these neuronal cell lines was investigated to determine if responsive cells had a consistent receptor profile.

In one experiment, cells from the SH-SY5Y neuroblastoma cell line were plated at  $2x10^5$  cells/ml/well in 12-well tissue culture plates. After one day in culture cells were either left untreated (Cn) or stimulated with 50 ng/ml artemin (ART) or 10  $\mu$ M retinoic acid (RA), a known inducer of differentiation of SH-SY5Y cells. Three days after factor addition, differentiation was assessed and cells were photographed. As shown in Figs. 8A-8C, artemin and retinoic acid induced strong differentiation responses in the SH-SY5Y cell line as indicated by their robust neurite outgrowth and neuronal morphology.

In another experiment, the ability of artemin to induce proliferation of the NBL-S neuroblastoma cell line was examined. NBL-S cells were plated at  $5x10^4$  cells/well on 48-well plates in standard medium, either without treatment or in the presence of 50 ng/mL of GDNF, artemin, or persephin. Actively dividing cells undergoing DNA synthesis 30 hrs after factor addition were detected using the BrdU (colorimetric) cell proliferation assay kit according to the manufacturer's instructions (Boehringer-Mannheim) and the data are shown in Fig. 8D. Like GDNF, artemin stimulated proliferation of NBL-S cells, whereas persephin did not.

To confirm that RET was a required component of the artemin receptor, the responsiveness of the neuroblastoma cell lines CHP126, CHP134, and SAN, which do not express RET, were examined. These cell lines did not respond to GDNF, neurturin or artemin (data not shown). Because both the SH-SY5Y and NBL-S cell lines express GFR $\alpha$ 1, GFR $\alpha$ 2, and GFR $\alpha$ 3, as well as RET, these experiments did not suggest the usage of a particular coreceptor by artemin.

#### Example 6

This example illustrates that artemin supports the survival of dopaminergic neurons.

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As shown above, artemin expression was not observed in the brain or spinal cord of embryonic rats at the age examined and was not observed in the fetal human brain, but was present at very low levels in some adult brain regions. Therefore artemin's ability to influence dopaminergic neurons from the rat embryonic ventral midbrain was compared to the survival promoting activity of the other GDNF ligand family members.

Embryonic day 14 ventral mesencephalon cultures were prepared by removing the entire mesencephalon into cold Leibovitz's L15 + 6 mg/ml glucose and dissecting the tissue while keeping it on ice. Following dissection, the tissue was digested in a mixture of dispase (1 mg/ml, Sigma) and collagenase (1 mg/ml, Worthington Biochemical) for 25 min. Tissue was then washed twice with modified N2 media and triturated 35 times. Cell density and viability was assessed using a hemocytometer to count trypan blue excluding cells. Cells were plated at 20,000 cells/well on 8-well chamber slides (coated with 125 ng/ml poly-D-lysine and 25 ng/ml laminin) in serum-free medium consisting of DME/Hams F12 (1:1), 1 mg/ml BSA, 5 μM insulin, 10 nM progesterone, 100 μM putrescine, 30 nM selenium, 10 ng/ml rat transferrin, 100 U/ml penicillin, 100 U/ml streptomycin. Factors were added within 15 min of plating. After 3 days in culture, the cells were fixed, stained for tyrosine hydroxylase, and the number of tyrosine hydroxylase staining (TH+) neurons were counted. The results from two independent experiments are shown in Fig. 7F as the percentage of surviving TH+ neurons over control.

Interestingly, artemin supported the survival of dopaminergic midbrain neurons, although there is no apparent artemin expression in the ventral midbrain at this age. Therefore, artemin can promote survival of central as well as peripheral neurons, and indicates that receptors for artemin are present in both these populations.

In summary, all of the biological responses of artemin in peripheral and central neurons, and in neuronal cell lines, suggest that artemin utilizes receptor components similar to or overlapping with GDNF and neurturin.

#### Example 7

This example illustrates that artemin signals through RET using the former orphan receptor GFRα3 as a coreceptor.

Because artemin, GDNF and neurturin exhibit similar activity profiles for peripheral and central neuronal populations and because artemin's activity correlated with RET expression in neuroblastoma cell lines, artemin's ability to activate RET in primary cultured SCG neurons and in the artemin-responsive neuroblastoma cell line NBL-S was examined by performing Western blot analysis of Ret phosphorylation and MAP kinase activation as described previously (Baloh et al., 1997, supra; Creedon et al., 1997, supra).

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England Biolabs).

In brief, SCG neurons were dissected from postnatal day 1 (P1) rats and maintained in NGF-containing medium for 5 days, after which they were deprived of NGF by switching to NGF-free medium in the presence of anti-NGF antibodies. After 2 hours without NGF, neurons were switched to medium containing NGF, GDNF or artemin at 50 ng/ml for 20 minutes. Cells were washed in cold PBS, and 15 collected in immunoprecipitation buffer (1 mM EDTA/1 mM EGTA/0.2mM NaVO<sub>3</sub>/1 mM Pefabloc/1 uM pepstatin A/10ug/ml leupeptin/2 ug/ml aprotinin/1% Triton X-100/0.5% Nonidet P-40/150 mM NaCl in 10 mM Tris, pH 7.4). The lysates were incubated with 30 µl agarose-conjugated anti-phosphotyrosine antibodies (Calbiochem) at 4°C for 1 hr. Beads were then washed three times with 20 immunoprecipitation buffer, resuspended in SDS sample buffer, and boiled for 5 min. Samples were separated using SDS-PAGE, and transferred to nitrocellulose membranes. Membranes were blocked for 1 hr at 25°C in TBS containing 5% dry milk, incubated overnight 4°C with a 1:300 dilution of anti-Ret antibody (C-19; Santa Cruz), then washed (3X in TBS containing 0.1% Tween-20) and incubated in a 25 1:10,000 dilution of anti-rabbit antibodies conjugated to HRP (Jackson Immunoresearch). Membranes were washed 3X, incubated with SuperSignal ULTRA (Pierce) for 5 minutes, then exposed to film. For MAP kinase (MAPK) assays, a portion of the total lysate wars removed before immunoprecipitation, separated by SDS-PAGE, and immunoblotted using an anti-phospho-MAP-kinase antibody (New 30

NBL-S cells were plated at  $3x10^5$  cells/well in 6-well plates, and after 46 hr switched to low-serum (0.5%) medium for 2 hr, and then stimulated with NGF, GDNF or artemin (50 ng/ml) for 20 min. Cells were then collected and analyzed as above. For PI-PLC treatment, a parallel set of NBL-S cells were plated as above but treated with 500 mU/ml of PI-PLC (Boehringer-Mannheim) for 1 hr, and washed with PBS and stimulated with factors as above.

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The results are shown in Figs. 9A and 9B. Like GDNF, artemin induced tyrosine-phosphorylation of RET in SCG neurons and in the NBL-S cell line, and this activation was effective in eliciting downstream signaling as measured by activation of the MAP kinase pathway. Furthermore, pretreatment of the NBLS cell line with the enzyme PI-PLC, which specifically cleaves GPI-anchored proteins from the cell surface, abolished artemin's ability to activate RET and all downstream signaling, but as expected had no effect on NGF signaling (Figure 9B). Therefore, these data indicate that like the other members of the GDNF ligand family, artemin signals through the RET receptor tyrosine kinase, and requires a GPI-anchored coreceptor to do so.

To investigate which if any of the known coreceptor(s) artemin can utilize to activate RET, its ability to directly bind to soluble Fc-fusion forms of GFRa1, GFRα2, or GFRα3 was assessed. GFRα1-Fc, GFRα2-Fc and GFRα3-Fc fusion proteins were obtained from R&D Systems. Binding assays were performed similarly to assays previously described (Sanicola et al., Proc. Natl. Acad. Sci., USA 94:6238-6243, 1997). Recombinant GDNF, neurturin, artemin, persephin, or purified bovine serum albumin (BSA; Pierce) were coated to Nunc-Immuno MaxiSorp microtiter plates at 325 ng/ml in TBS (10mM Tris-HCl pH 7.5, 150 mM NaCl) for 1 hour at 25°C. Plates were washed (3X in TBS/0.03% Tween-20), and then blocked (blocking solution: TBS/1% BSA) for 1 hr at 25°C. Receptor bodies diluted in blocking solution were added and incubated for 2 hr at 25°C, washed 5X, then incubated with 1:10,000 dilution of HRP-conjugated anti-human Fc antibodies in blocking solution (Jackson Immunoresearch) for 45 minutes at 25°C. Finally, wells were washed 5X and the presence of HRP was assayed by addition of the chromogenic substrate 3,3', 5,5-tetramethylbenzidine for 5 min. The reaction was stopped by adding an equal volume of 0.5M H<sub>2</sub>SO<sub>4</sub>, and color was measured in a plate reader at 450 nm. Non-

specific binding of receptor bodies to the plates was measured in BSA coated wells, and this binding was subtracted from all other measurements. All experiments were performed in duplicate and the results are shown in Figs. 9C-9E.

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In this assay, GFR $\alpha$ 1-Fc was able to bind to both GDNF and neurturin with similar affinity, but not to persephin or artemin. GFR $\alpha$ 2-Fc bound to neurturin but not to GDNF or artemin, consistent with previous reports that GDNF can only bind to GFR $\alpha$ 2 in the presence of RET (Sanicola et al., 1997). Interestingly, the former orphan receptor GFR $\alpha$ 3-Fc was able to bind to artemin, but not to any of the other GDNF ligand family members, consistent with the inability of GFR $\alpha$ 3 and RET to form a functional receptor complex for GDNF, neurturin or persephin (Baloh et al., 1998, *supra*; Worby et al., 1998, *supra*). Each of the receptor bodies bound to their respective ligands with an apparent Kd of ~3nM, which is similar to the previously reported affinity between GFR $\alpha$ 1-Fc and GDNF in this assay (Sanicola et al., 1997). Therefore, artemin can bind to the former orphan receptor GFR $\alpha$ 3, with an affinity similar to GDNF and NTN binding to their respective receptors.

While the binding data above suggest that GFRα3/RET is the functional receptor for artemin, such direct binding studies alone have been unreliable in predicting all GDNF family receptor interactions because of the observation that RET can modulate receptor binding (Sanicola et al., 1997, *supra*). Thus, to directly test which receptor combinations can form functional receptors for artemin, expression plasmids for the individual GFRα coreceptors together with the Gal4-Elk/Gal4-Luc reporter system were transiently transformed into fibroblasts that stably express RET. This system, which utilizes the ability of the Gal4-Elk fusion protein to respond to MAP kinase activity and activate transcription of the Gal4-luciferase reporter, has been used previously to monitor NGF/TrkA activation of the MAP kinase pathway in PC12 cells (Vossler et al., *Cell* 89:73-82, 1997; York et al., *Nature* 392:622-626, 1998), and GDNF/RET activation of MAP kinase in neuroblastoma cell lines (Worby et al., 1998, *supra*; Worby et al., *J. Biol. Chem*.271:23619-23622, 1996).

Generation of the expression plasmids for rat GFR $\alpha$ 1 and human GFR $\alpha$ 2, as well as MG87 fibroblasts stably expressing human RET (RET-3T3), were described in detail previously (Baloh et al., 1997, *supra*; Creedon et al., 1997, *supra*). For generation of the GFR $\alpha$ 3 expression plasmid, a fragment containing the complete

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coding region of human GFR $\alpha$ 3 was PCR-amplified from a human pituitary cDNA library, and this fragment was directly cloned into the KpnI and BamHI sites of pCB6 (Brewer, *Methods in Cell Biol.* 43:233-245, 1994) using sites engineered into the PCR primers. The GAL4-Elk1 chimera expression plasmid was a gift of P. Stork (Vollum Institute, Oregon Health Sciences University).

NLF neuroblastoma cells, which naturally express GFRα2/RET (Tansey et al., submitted), or the RET-3T3 cells were plated at 85,000 cells/well in 12-well plates, and cotransfected with the reporter plasmids Gal4-Luc and CMV-Gal4-Elk (250 ng/well and 50 ng/well, respectively), CMV-lacZ (50 ng/well) for transfection normalization, one of the CMV-GFR\alpha expression plasmids (500 ng/well), and pBluescript (650 ng/well) as carrier for a total of 1.5µg DNA/well using the Superfect reagent (Qiagen) according to the manufacturer's instructions. RET-3T3 fibroblasts were exposed to DNA/Superfect mixtures at 37°C overnight, washed and placed in low-serum (0.5%) medium, and stimulated with 50 ng/ml GDNF, artemin or persephin for 6-8 hours before collection (48 hours post-transformation). NLF neuroblastoma cells were exposed to DNA/Superfect mixtures for 2 hours, placed in full-serum medium overnight for recovery, and then switched to low-serum (0.5%) medium containing 50 ng/m of GDNF, artemin or persephin for 24 hours before collection (48 hours post-transformation). Measurement of luciferase and β-galactosidase activity was measured using a luminometer and performed as described (Syaren et al., EMBO J 17:6010-6019, 1998). The average luciferase activity of duplicate samples was normalized to β-galactosidase activity of the cotransformed lacZ reporter, and fold activation was calculated by dividing the normalized activity in factor-treated cells by that in the no treatment control.

As seen in Fig. 10A, consistent with previous reports in multiple systems, GDNF activated RET signaling in cells expressing GFR $\alpha$ 1/RET or GFR $\alpha$ 2/RET, but not GFR $\alpha$ 3/RET (Baloh et al., 1998; Baloh et al., 1997; Jing et al., 1997; Sanicola et al., 1997; Suvanto et al., 1997; Worby et al., 1998). As predicted from the binding data, artemin was able to activate GFR $\alpha$ 3/RET, but not GFR $\alpha$ 2/RET receptor complexes. Interestingly, artemin was also able to activate RET in cells expressing GFR $\alpha$ 1/RET, although to a lesser degree than GDNF.

WO 00/18799 PCT/US99/22604

62

These results were confirmed in the NLF cell line, which is a more neuronal cell line that natively responds to GDNF by RET and MAP kinase activation (unpublished data). As shown in Fig. 10B, NLF cells transformed with GFR $\alpha$ 1 responded more intensely to GDNF, and became responsive to artemin, confirming the results observed in fibroblasts. Furthermore, transformation of NLF cells with GFR $\alpha$ 3, but not GFR $\alpha$ 2, allowed the cells to respond to artemin stimulation. Transformation of NFL cells with GFR $\alpha$ 3 also decreased the ability of the cells to respond to GDNF, presumably because CMV-driven overexpression of GFR $\alpha$ 3 decreases the relative amount of GFR $\alpha$ 2/RET complexes on the cells, thereby decreasing the number of functional GDNF receptors.

In summary, direct binding data and *in vitro* receptor activation experiments together indicate that artemin is the only known GDNF family ligand for the  $GFR\alpha 3/RET$  receptor complex, but like GDNF and neurturin it can also activate the  $GFR\alpha 1/RET$  receptor complex.

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### Example 8

This example illustrates the specificity and cross-talk in interactions of members of the GDNF ligand and  $GFR\alpha$  receptor families.

A schematic interaction diagram of the GDNF ligand family members and GFRα receptor family members is presented in Fig. 11. Similar to many other ligand/receptor systems, including the neurotrophins, the system is characterized by having preferred ligand/receptor pairs, however cross-talk between the different ligands and receptors is apparent. The addition of artemin to this diagram reveals several new features. First, both direct receptor binding experiments and receptor activation experiments presented herein indicate that artemin is the only GDNF family ligand capable of utilizing the GFRα3/RET receptor complex. One additional study indicated that GDNF was able bind to GFRα3 in the presence of RET (Trupp et al., 1998), however this interaction was of low-affinity in that it required cross-linking to be observed, and its relevance is unclear because it does not lead to RET activation in multiple experimental paradigms *in vitro* (Baloh et al., 1998; Trupp et al., 1998; Worby et al., 1998). Second, the GFRα1/RET receptor complex is highly

WO 00/18799 PCT/US99/22604

63

promiscuous, and although results from most in vitro paradigms agree that it is the preferred receptor for GDNF, it is believed that neurturin and artemin also utilize the GFRα1/RET receptor complex in some systems *in vivo*. Recent observations regarding differences in peripheral neuron losses between GDNF and GFRα1-deficient mice suggest that GDNF can use GFRα2/RET as a receptor *in vivo*, and confirm that alternative ligand/receptor interactions may have biological importance in the GDNF family (Cacalano et al., 1998; Enomoto et al., 1998). Furthermore, a recent paper analyzing neurotrophin knockout mice concluded that NT-3 can signal through TrkB *in vivo*, an interaction observed many years earlier *in vitro* but was thought to be irrelevant *in vivo* (Farinas I et al., 1998; Ip et al., 1993). Therefore, consideration of all possible ligand/receptor interactions identified *in vitro* is often necessary to understand results of *in vivo* analysis of neurotrophic factor influences.

Results from direct binding and RET activation experiments presented herein further suggest that like the GDNF-GFR $\alpha$ 2 interaction, the artemin-GFR $\alpha$ 1 interaction appears dependent on the presence of RET, as direct binding of artemin to GFR $\alpha$ 1-Fc, or GDNF to GFR $\alpha$ 2-Fc receptor bodies was not observed (Fig. 9C; Sanicola et al., 1997). This may be due to the nature of the binding assay which utilizes soluble receptor bodies binding to immobilized ligand. However, this may alternatively reflect an additional level of specificity available to the GDNF ligand/GFR $\alpha$  system, in cases where the GFR $\alpha$  coreceptors are expressed in the absence of RET (Baloh et al., 1997; Golden et al., 1998; Trupp et al., 1997; Yu et al., 1998). In these situations (i.e. GFR $\alpha$ 1 in injured peripheral nerve, and GFR $\alpha$ 2 expression in cerebral cortex), where coreceptors are expressed in "trans" and are hypothesized to secrete or present ligand/coreceptor complexes to cells or axons expressing RET, only the RET-independent subset of binding interactions would be possible.

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<u>Deposit of Plasmid</u>: The following plasmid has been deposited under the terms of the Budapest Treaty, with the American Type Culture Collection, 10801 University Blvd., Manassas, VA 20110-2209. The accession number indicated was assigned after successful verification of the presence of the plasmid in the deposit, and the requisite fees have been paid. Access to said plasmid will be available during pendency of the patent application to one determined by the Commissioner to be

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entitled thereto under 37 CFR 1.14 and 35 USC 122. All restriction on availability of said plasmid to the public will be irrevocably removed upon the granting of a patent based upon the application. Moreover, the designated deposits will be maintained for a period of thirty (30) years from the date of deposit, or for five (5) years after the last request for the deposit, or for the enforceable life of the U.S. patent, whichever is longer. Should the plasmid become nonviable or be inadvertently destroyed, it will be replaced with a viable plasmid. The deposited materials mentioned herein are intended for convenience only, and are not required to practice the present invention in view of the description herein, and in addition, these materials are incorporated herein by reference.

	Name of Plasmid	Deposit Date	ATCC No.
15	phART	December 22, 1998	

In view of the above, it will be seen that the several advantages of the invention are achieved and other advantageous results attained.

As various changes could be made in the above methods and compositions without departing from the scope of the invention, it is intended that all matter contained in the above description and shown in the accompanying drawings shall be interpreted as illustrative and not in a limiting sense.

All references cited in this specification, including patents and patent applications, are hereby incorporated by reference. The discussion of references herein is intended merely to summarize the assertions made by their authors and no admission is made that any reference constitutes prior art. Applicants reserve the right to challenge the accuracy and pertinency of the cited references.

PCT/US99/22604

### What is Claimed is:

- 1. An isolated and purified growth factor comprising an artemin amino acid sequence or a conservatively substituted variant thereof or a fragment thereof of at least 8 contiguous amino acids.
- 2. The isolated and purified growth factor of claim 1 which promotes survival of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosine-hydroxylase-expressing dopaminergic ventral midbrain neurons.
- 3. The isolated and purified growth factor of claim 1 comprising a mammalian sequence which is at least 75% identical to SEQ ID NO:19, SEQ ID NO:33 or a conservatively substituted variant thereof.
- 4. The isolated and purified growth factor of claim 3 comprising a human polypeptide sequence as set forth in SEQ ID NO: 3, SEQ ID NO: 4, SEQ ID NO:5.
- 5. The isolated and purified growth factor of claim 4 comprising a human pro-artemin as set forth in SEQ ID NO:40 or a human pre-pro-artemin as set forth in SEQ ID NO:26 or SEQ ID NO:32 or a conservatively substituted variant thereof or a polypeptide comprising a non-artemin pre-pro- region and the human polypeptide sequence.
- 6. The isolated and purified growth factor of claim 3 comprising a mouse polypeptide sequence as set forth in SEQ ID NO:34, SEQ ID NO:35, SEQ ID NO:36 or a conservatively substituted variant thereof.
- 7. The isolated and purified growth factor of claim 6 comprising a mouse pro-artemin as set forth in SEQ ID NO:41 or a mouse pre-pro-artemin as set forth in SEQ ID NO:27 or a conservatively substituted variant thereof or a polypeptide comprising a non-artemin pre-pro- region and the mouse polypeptide.

- 8. The isolated and purified growth factor of claim 1 comprising an amino acid sequence identical to the artemin polypeptide encoded by the human cDNA contained in the DNA deposited with ATCC on December 22, 1998.
- 9. The isolated and purified growth factor of claim 1 comprising an artemin polypeptide produced by a process comprising the steps of:
- (a) transforming a host cell with human artemin cDNA clone deposited with ATCC on December 22, 1998 operably linked to expression regulatory elements and
  - (b) expressing artemin polypeptide encoded by the clone.
  - 10. An isolated and purified polypeptide comprising:

- (a) a pre- region of human artemin as set forth in SEQ ID NO:48 or a preregion of mouse artemin as set forth in SEQ ID NO:49,
- (b) a pro- region of human artemin as set forth in SEQ ID NO:50 or a proregion of mouse artemin as set forth in SEQ ID NO:51,
- (c) a pre-pro- region of human artemin as set forth in SEQ ID NO:52 or a pre-pro- region of mouse artemin as set forth in SEQ ID NO:53, or
  - (d) a conservatively substituted variant of (a), (b) or (c)...
- 11. A pan-growth factor comprising the artemin polypeptide fragment of claim 1 and a fragment of at least one other growth factor from the TGF- $\beta$  superfamily.
- 12. A nucleic acid comprising a polynucleotide encoding the pan-growth factor of claim 11.
- 13. A composition comprising the growth factor of claim 1 and a GFR $\alpha$  polypeptide.
- 14. The composition of claim 13 wherein the GFR $\alpha$  polypeptide is a GFR $\alpha$ 3 polypeptide or a GFR $\alpha$ 1 polypeptide.

- 15. An isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto comprising a nucleotide sequence encoding a growth factor of claim 1 or a fragment of said nucleotide sequence consisting of at least 15 contiguous nucleotides.
- 16. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 15 comprising a nucleotide sequence encoding an artemin polypeptide which promotes survival of trigeminal ganglion neurons, nodose ganglion neurons, superior cervical ganglion neurons, and tyrosine-hydroxylase-expressing dopaminergic ventral midbrain neurons wherein said nucleic acid molecule specifically hybridizes to a mature human artemin nucleotide sequence as set forth in SEQ ID NO:6, SEQ ID NO:7 or SEQ ID NO:8 or to a mature mouse artemin nucleotide sequence as set forth in SEQ ID NO:37, SEQ ID NO:38 or SEQ ID NO:39.
  - 17. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 16 comprising a nucleotide sequence encoding an artemin polypeptide as set forth in SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5, SEQ ID NO:34, SEQ ID NO:35 or SEQ ID NO:36.
  - 18. The isolated and purified nucleic acid molecule or nucleic acid molecule complementary thereto of claim 17 comprising a nucleotide sequence as set forth in SEQ ID NO:6, SEQ ID NO:7, SEQ ID NO:8, SEQ ID NO:37, SEQ ID NO:38, SEQ ID NO:39 or SEQ ID NO:44.
  - 19. A vector comprising expression regulatory elements operably linked to the nucleic acid molecule of claim 15.
    - 20. A host cell transformed with the vector of claim 19.
  - 21. The isolated and purified nucleic acid molecule of claim 15 comprising ATCC deposit made on December 22, 1998.

22. A host cell transformed with the vector of claim 21.

WO 00/18799

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- 23. The isolated and purified nucleic acid molecule of claim 15 comprising a polynucleotide encoding a polypeptide selected from the group consisting of a human pro-artemin as set forth in SEQ ID NO:41, a human pre-pro artemin as set forth in SEQ ID NO:26 or SEQ ID NO:32, a mouse pro-artemin as set forth in SEQ ID NO:42, a mouse pre-pro-artemin as set forth in SEQ ID NO:29 and a polypeptide comprising a non-artemin pre-pro- region sequence and a human or mouse mature artemin amino acid sequence.
  - 24. The isolated and purified nucleic acid molecule of claim 23 comprising a human pro-artemin nucleotide as set forth in SEQ ID NO:42, a human pre-pro-artemin nucleotide as set forth in SEQ ID NO:24, SEQ ID NO:30 or SEQ ID NO:44, a mouse pro-artemin nucleotide as set forth in SEQ ID NO:43, or a mouse pre-pro-artemin nucleotide as set forth in SEQ ID NO:27.
  - 25. A recombinant nucleic acid molecule comprising an artemin nucleotide sequence or complement thereto wherein the artemin nucleotide sequence encodes an amino acid sequence selected from the group consisting of a pre-pro-artemin polypeptide, a pro-artemin polypeptide, a mature artemin polypeptide, a conservatively substituted variant thereof and a fragment thereof having at least 8 contiguous amino acids.
  - 26. The isolated and purified polynucleotide of claim 15 which is an artemin antisense oligonucleotide.
  - 27. An isolated and purified nucleic acid molecule comprising a polynucleotide encoding:
  - (a) a pre- region of artemin as set forth in SEQ ID NO:54 or SEQ ID NO:55;
- 5 (b) a pro- region of artemin as set forth in SEQ ID NO:56 or SEQ ID NO:57;

WO 00/18799 PCT/US99/22604

- (c) a pre-pro- region of artemin as set forth in SEQ ID NO:58 or SEQ ID NO:59; or
  - (d) a conservatively substituted variant of (a), (b) or (c).
- 28. An isolated and purified antibody which specifically reacts with the artemin polypeptide or fragment of claim 1.
- 29. A method for detecting expression of an artemin polypeptide in a sample comprising contacting the sample with an antibody according to claim 28 and detecting binding of the antibody to the artemin polypeptide.
- 30. A method for detecting expression of an artemin mRNA in a sample which comprises detecting a polynucleotide in the sample that specifically hybridizes to a polynucleotide consisting of SEQ ID NO:9.
- 31. The method of claim 30, wherein detecting the polynucleotide comprises:
- (a) contacting mRNA of the sample with a polynucleotide that specifically hybridizes to a polynucleotide consisting of SEQ ID NO:6; and
- (b) detecting the existence of a hybridization complex between the polynucleotide and the artemin mRNA.
  - 32. The method of claim 30, wherein the detecting step comprises:
- (a) producing a cDNA from the artemin mRNA using the reverse transcription method;
- (b) contacting the cDNA with at least two oligonucleotides that specifically hybridize to the cDNA to define a region of the cDNA to be amplified;
  - (c) amplifying the cDNA region; and

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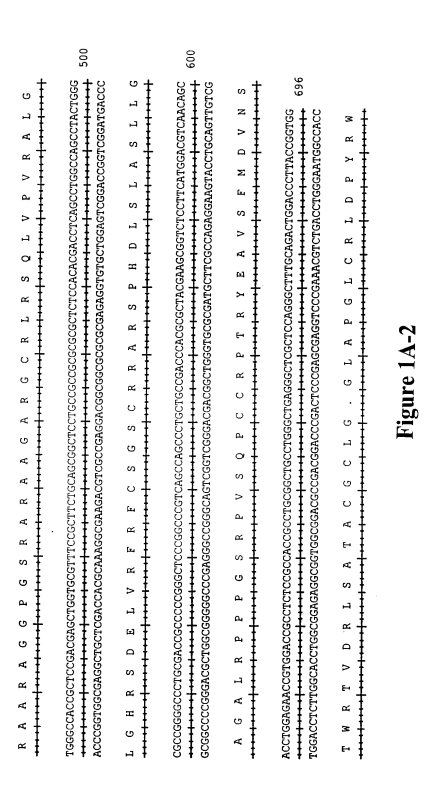
(d) detecting the amplified cDNA region.

- 33. A method for providing trophic support to and/or for producing differentiation of a cell comprising treating the cell with an effective amount of an artemin polypeptide or fragment thereof.
- 34. The method of claim 33, wherein the treating step comprises administering to the cell the artemin polypeptide or fragment with or without a  $GFR\alpha3$  polypeptide.
- 35. The method of claim 33, wherein the target cell is within a patient and the treating step comprises administering to the patient the artemin polypeptide or fragment with or without a GFR $\alpha$ 3 polypeptide.
- 36. The method of claim 33, wherein the target cell is within a patient and the treating step comprises administering to the patient a polynucleotide encoding the artemin polypeptide or fragment.
- 37. The method of claim 33, wherein the artemin polypeptide is expressed by a cell implanted into the patient.
- 38. The method of claim 33, wherein the target cell is a neuron in a patient suffering from peripheral neuropathy, amyotrophic lateral sclerosis, Alzheimer's disease, Parkinson's disease, Huntington's disease, ischemic stroke, acute brain injury, acute spinal chord injury, a nervous system tumor such as neuroblastomas, multiple sclerosis, infection or an enteric disease such as idiopathic constipation or constipation associated with Parkinson's disease, spinal cord injury or use of opiate pain-killers or the target cell is a non-neuronal cell in a patient suffering from small cell lung carcinoma.

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200	CCCCGCCGGCCACCTGCCGGGTAGGTGAGGGGGGGGGGG
	WPLWLC, AASQRPPWAPRPAALPPAKAPRLSWR
	acceccagacceagaccagcatcetccegaegeacceagegecegegegegegegegegegegeg
100	TGGCCGCTCTGGCTGAGCAGCGTCGCAGAGGCCTCCCTGGGCTCCGCGCCCCGCAGCCCTGCCCCCGCGAAGGCCCCCCGCGCTGTCCTGGCGTC

Figure 1A-1



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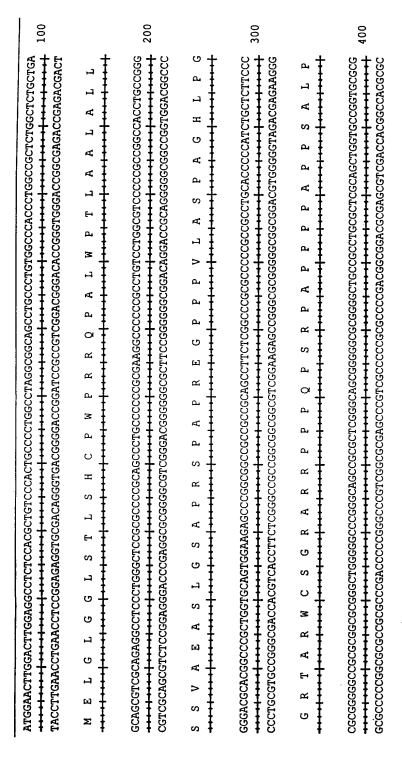
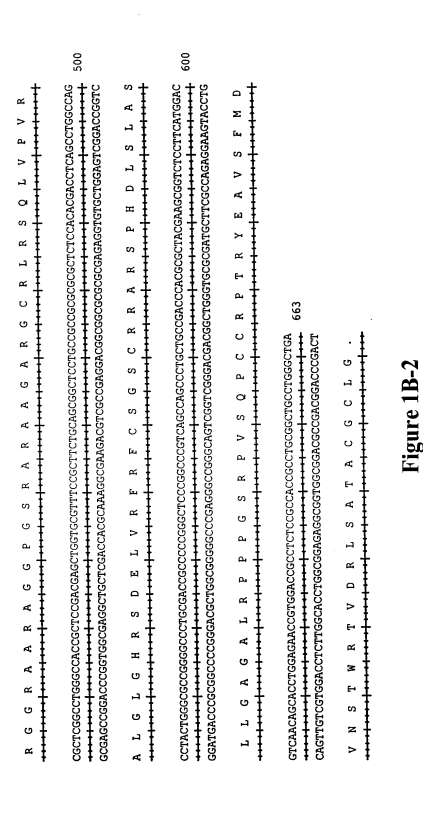


Figure 1B-1



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Figure 1C-1

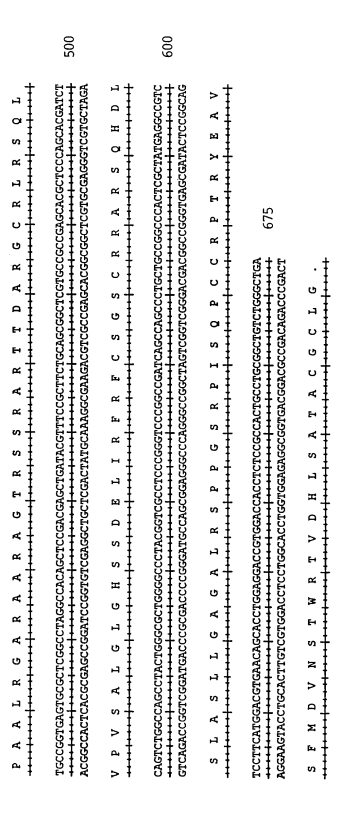


Figure 1C-2

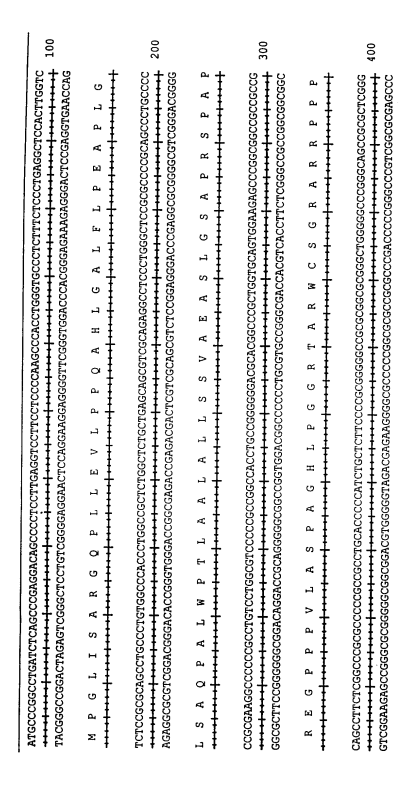
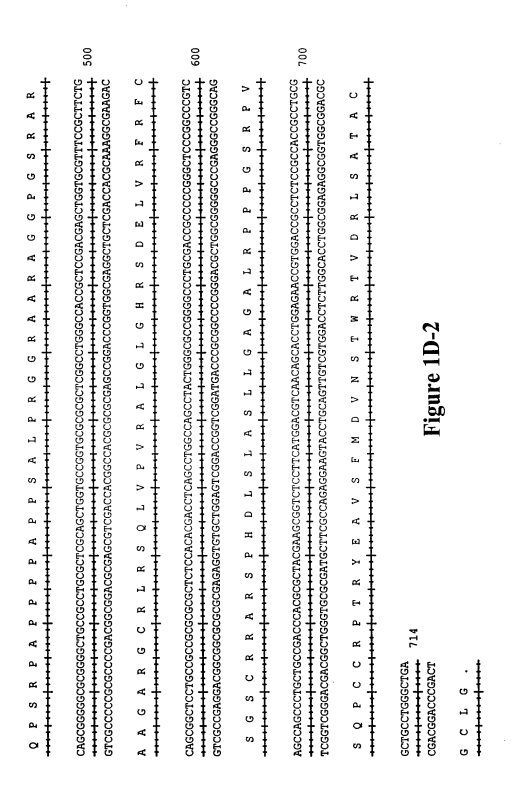


Figure 1D-1

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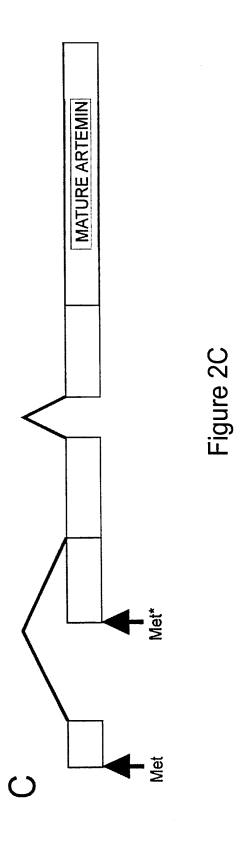
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Figure 2B



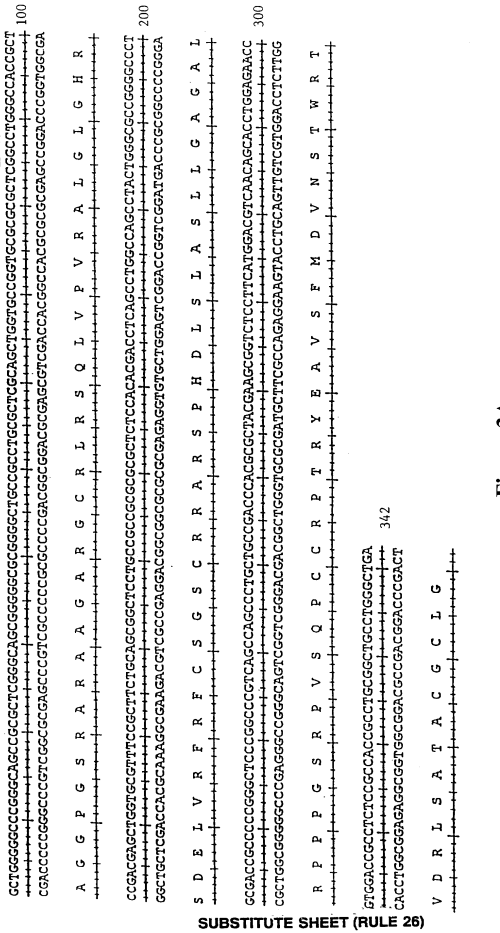
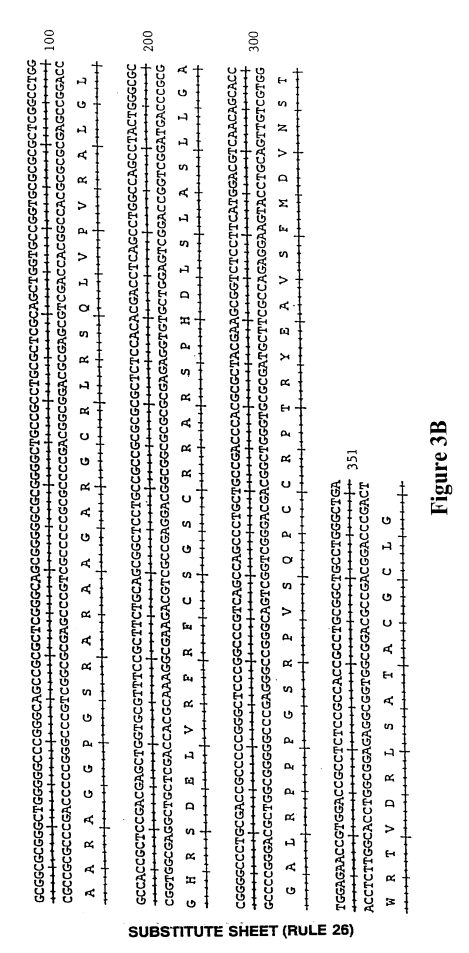
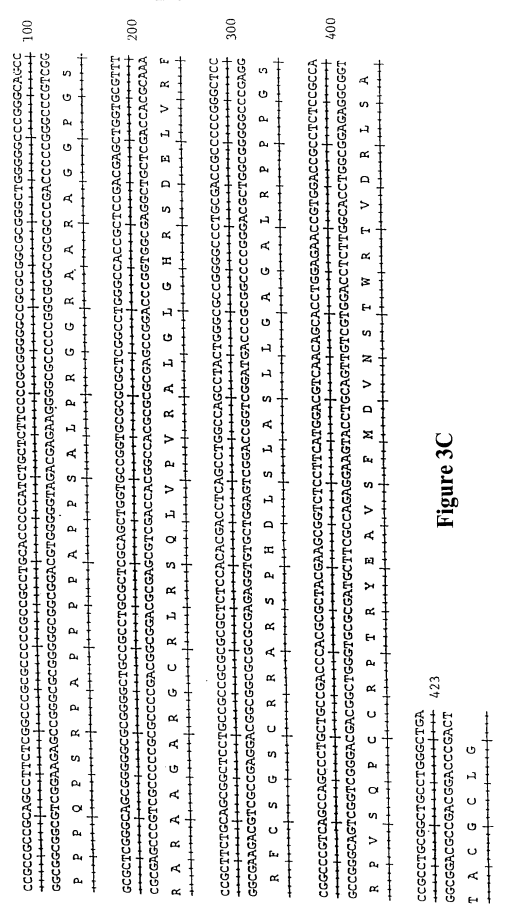


Figure 3A





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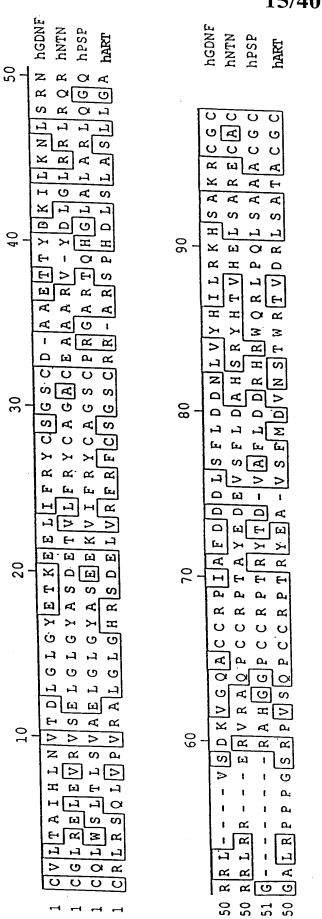
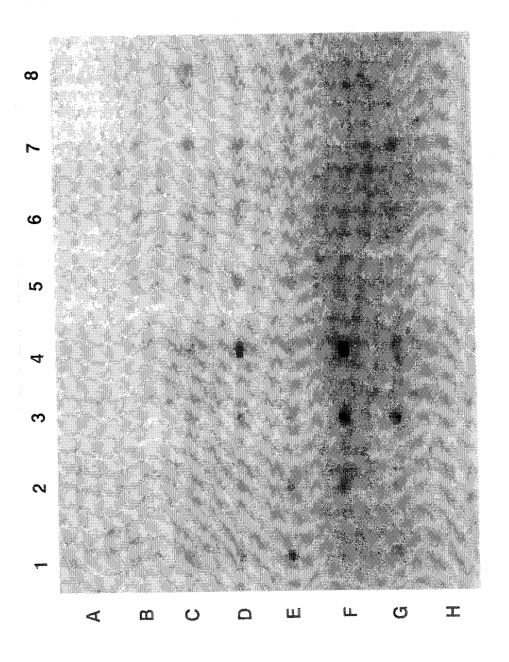


Figure 4

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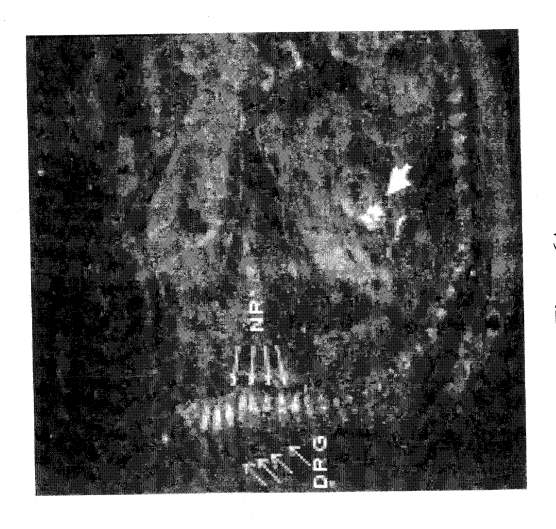
**Figure 5**A

WO 00/18799 PCT/US99/22604

	1	2	3	4	5	6	7	8
A	whole brain	amygdala	caudate nucleus	cerebellum	cerebral cortex	frontal lobe	hippocampus	medulla oblongata
В	occipital pole	putamen	substantia nigra	temporal lobe	thalamus	subthalamic nucleus	spinal cord	
C	heart	aorta	skeletal muscle	colon	bladder	uterus	prostate	stomach
D	testis	ovary	pancreas	pituitary gland	adrenal gland	thyroid gland	salivary gland	mammary gland
E	kidney	liver	small intestine	spleen	thymus	peripheral leukocyte	lymph node	bone marrow
F	appendix	lung	trachea	placenta				
G	fetal brain	fetal heart	fetal kidney	fetal liver	fetal spleen	fetal thymus	fetal lung	
H	yeast total RNA	yeast tRNA	E. coli rRNA	E. coli DNA	Poly r(A)	human C <sub>0</sub> t DNA	human DNA	human DNA

Figure 5B

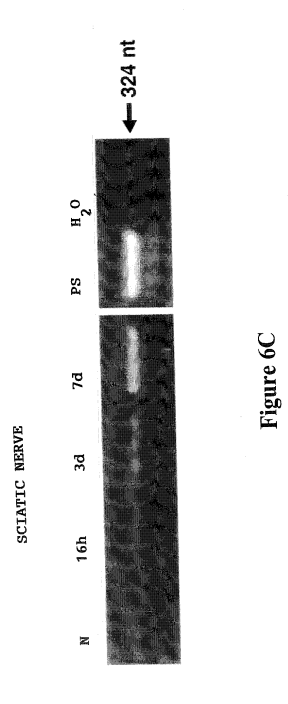
# Figure 6A

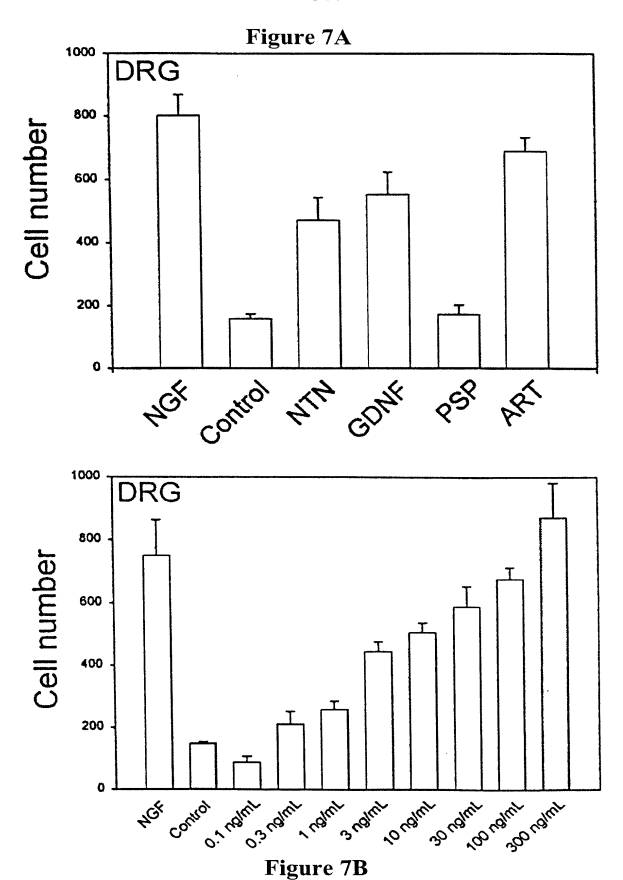


## Figure 6B



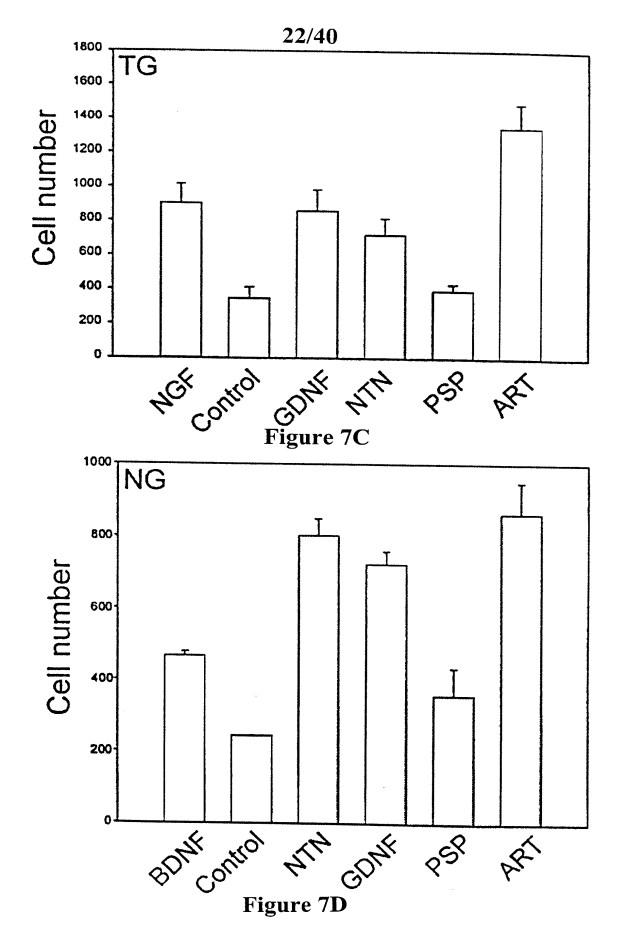
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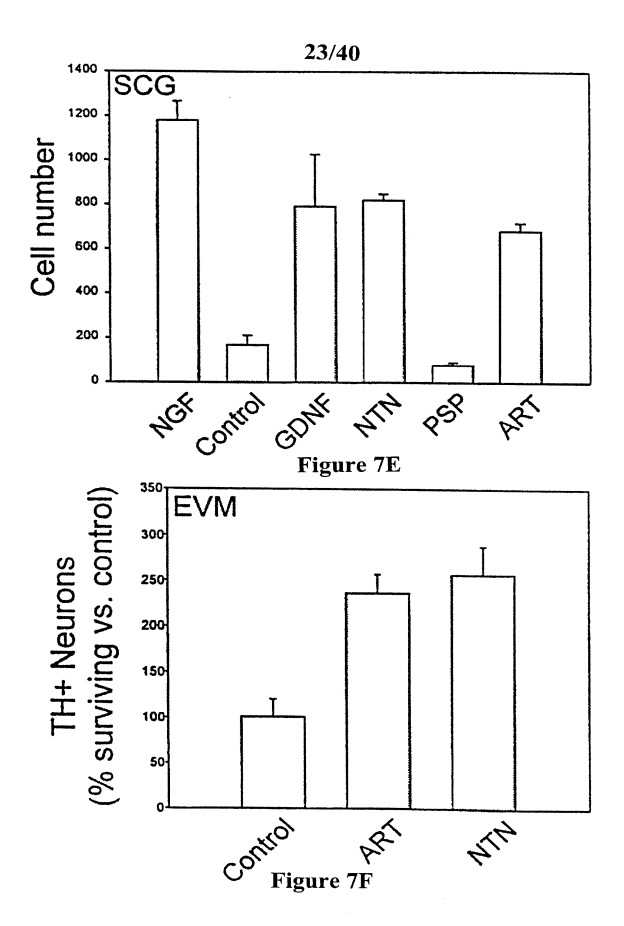


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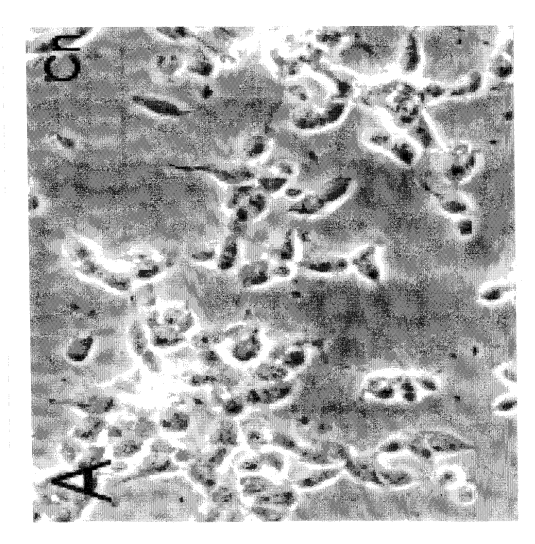


Figure 8A

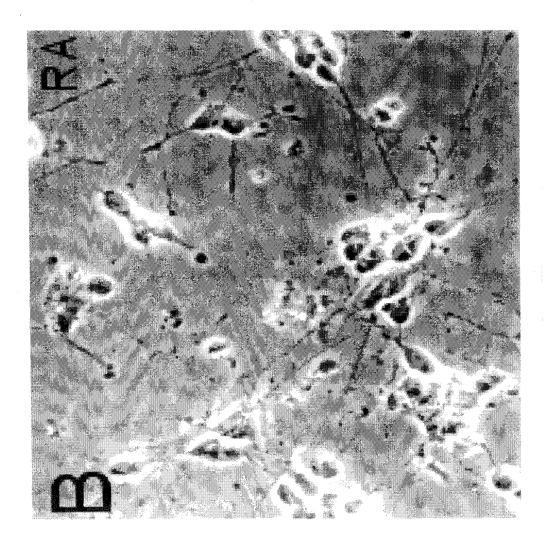
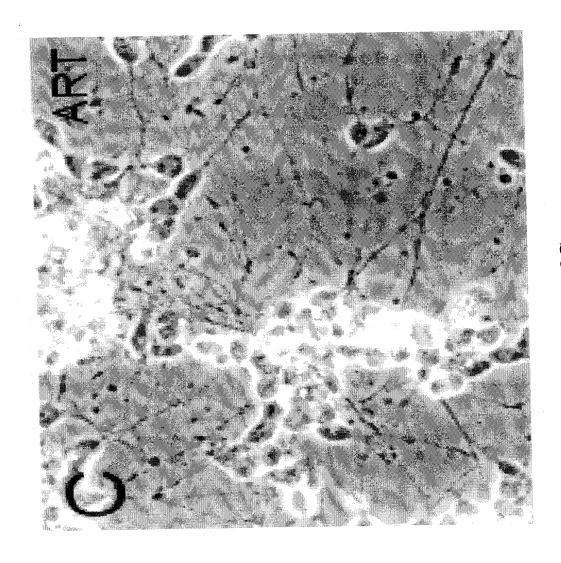


Figure 8B

## Figure &C



WO 00/18799 PCT/US99/22604



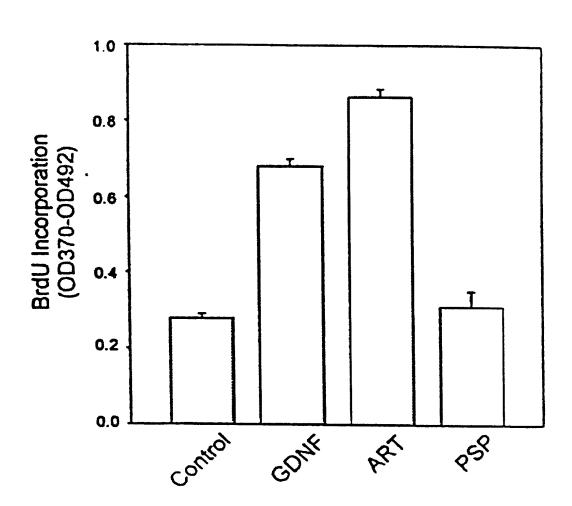
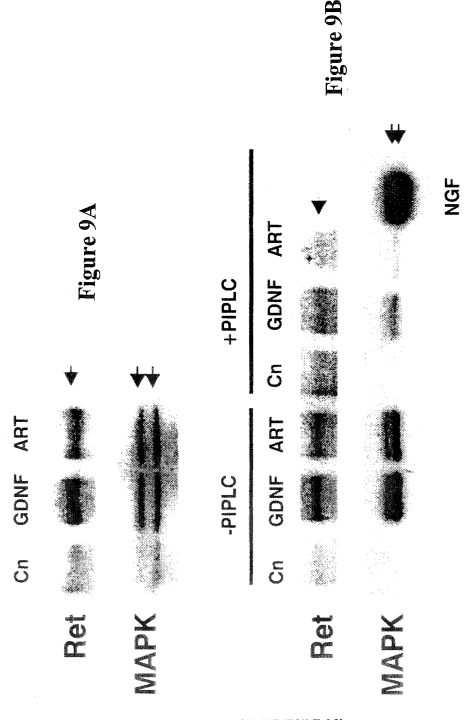
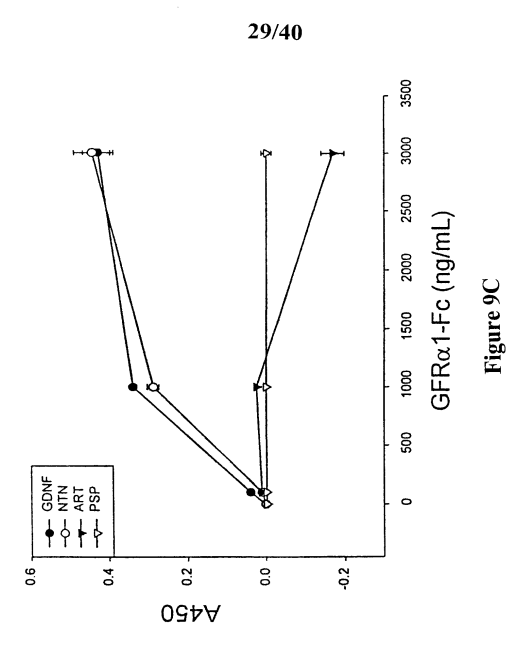


Figure 8D

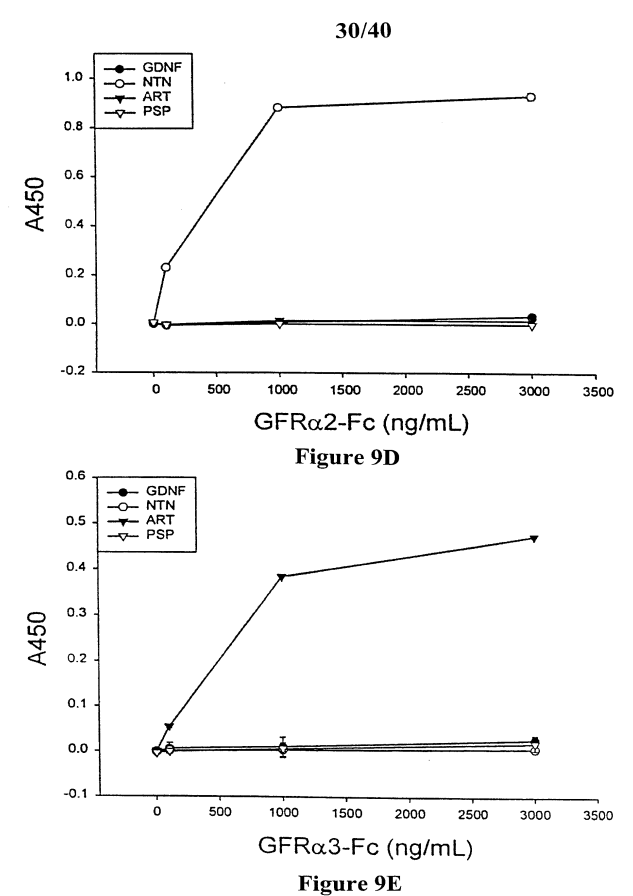


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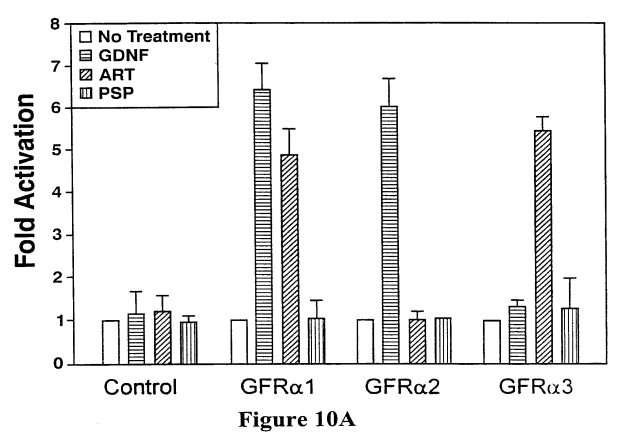


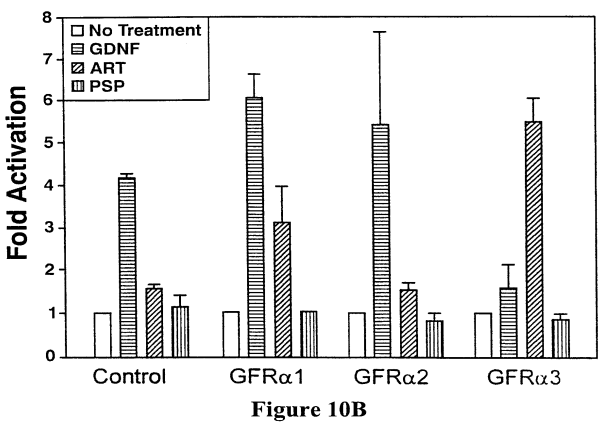
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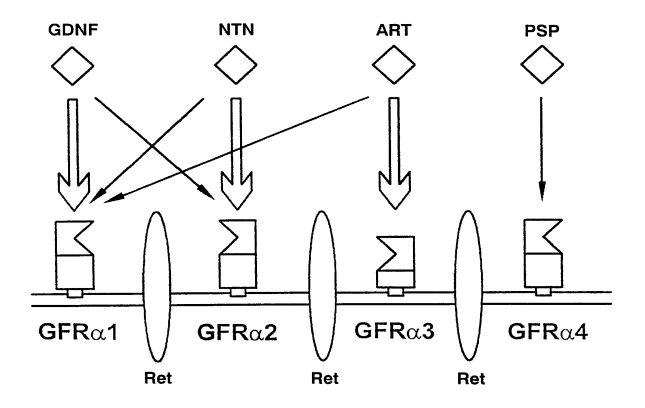


Figure 11

33/40  $\Sigma >$ > **S** A A X X 回回 ם M M S A O 2 2 니 도 SA H Н بدأ بدأ A H ഗ ഗ 田田 Σ> R R A S II U U بدأ بدأ |>>  $\alpha \alpha$ ШΩ > > o z 回回 > > HH E S D<sub>1</sub> S<sub>2</sub> ы ы Ø AS **E** шш ΩΩ P  $\vdash$ 卫民 म म HE 니 니 HH S 33 S HH HH A A H zz $\alpha \circ$ P P AB ΩΩ AA ×× U U **U U** Di S RA L L HЧ ቧ ቧ U U HHHH က က HO C C PS 4 ΩΩ X X z n **ს U U** BB OZ  $\vdash$   $\alpha$ H H HH ם HH 民民 33 G S C ΣΣ A A **>** 1 нн U U 0 A A нн K H ZZ H H HΩΩ H  $\alpha \circ$ A D SS K K > < Н H ΣΣ U U ഗ ഗ HH **S** S ା ପ  $\times \times$ 00 zzሷ ሷ တ တ मि मि ር ር  $\vdash$ ZI ഗ ഗ 00 $\alpha$ 工 工 니니 ᆸᅱ 7  $\times \times$ 田民 U U Н OK K Þ K 0 0 ł S ΣΣ Z Z ДΗ K K 田田 > > S ΣΣ ບ ບ S W D D KK च च HE SH II 저 저 머머 L K K ഗ ഗ **U U** म्य भ्य 00 HH 4 0 田田 工 工 33 υ U נבו נבו ZZ o民民 нн HO vው ው 4 4 > > ט ט ם ם ΣΣ L ΥX  $\Sigma \alpha$ XX 回回 KK > Σ ZX K K a a ı  $\overline{\mathcal{O}}$ S **5** ΩΩ L SS AA AA HH G HH ×× ZU R R > H **A A** SX I I > > A A םם S त्मि सि нн Σ I C C HH  $\vdash$ X X AA R R zz**A A** JΣ HA S 民民 U U U U дд 田田 Н > ם 교 SO HH ር ር 回 田 HH HH H> ΩZ **P** zz民 民 υ U × × ቧ ቧ ΣΣ L ם 44 民民 4 ΩΩ H H  $\Omega$ A A U U QЫ HH Д လ လ O O HH SA **P** HH HI  $\cup$   $\cup$ 00 > > X X L 百氏 0 0 ZZ ОШ P XX ΩΩ 0 ט ט U U H O H民民 民民 ZI A A Н 0 0 니 년 HH S ם R R 44 шш L AA R S <u>ს</u> ს म्य म्य Ø 44 ΞΞ Z 回回 ×× HH R HH 百公 ZΩ HB QQ r S HH UU H X K U & 4 0 I U U G G HH ध ध 日日 A A 田田 П 民工 OO ΩΩ H ΣΣ 디디 民民 A A H H A A A A > U S တတ AA U U ΗН ΣΟ hGFRa3 P mGFRa3 S hGFRa3 N mGFRa3 N hGFRa3 F. mGFRa3 E. hGFRa3 N mGFRa3 N hGFRa3 C mGFRa3 C hGFRα3 E | mGFRα3E ( mGFRa3M hGFRa3 M ĸ mGFRa3R hGFRa3 K mGFRa3 K hGFRa3 hGFRa3 hGFRa3

Figure 12

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## Figure 1.

Figure 14A-1

### 35/40

TAGTAAGGTTCCCAGTGCAGCTACTTCTGCTGGTTGAGTCTAGCTGTTGCTCTCTCT
TAGTAAGGTTCCCAGTGCAGCTACTTCTGCTGGGTT ATCATTCCAAGGGTCACGTCGATGAAGACGACCCAA  I V R F P V Q L L L L G  S Q C S Y F C W V  S K V P S A A T S A C T
** **** **** **** **** **** **** **** ****

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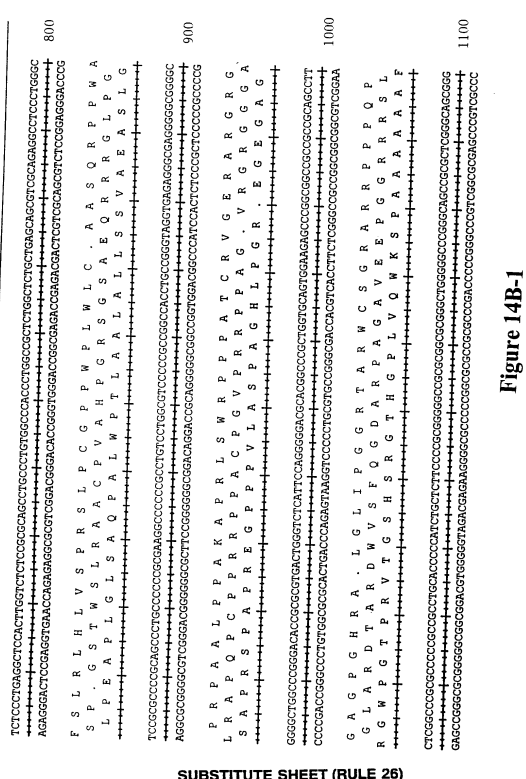
Figure 14A-2

36/40

	200		009		700	
GTWTWRPLHAVPLPLA. AAGEWFSQ. LLPGTEE ELGLGGGLSTLSHCPWPRRQVSGSPSDSYLVLRK WNLDLEASPRCPTAPGLGGR. VVLPVTPTWY. GK	GGCGGCTTGACTGGTGAGGGAGAGCAGGGCTTGGCTTGG	RRLDW. GRAGLGLGSG. VWEGKWSGRDQVNGRRS GGLTGEGEQGLAWAAVRCGRENGQGGTR. MGGG AA. LVRESRAWLGQRLGVGGKMVREGPGEWEEE	CGGGACTTCTCTGAATGGTCGGTGCACTCAGGTGATTCCTCCCCTGGGCTCCCAGAGGCAACCCATTATACTGGAACCTAGGCCCTTCCTGAGTTT 6	GTSLNGRCTQVIPPLGSQRQQTHYTGT. ALPEF AGLL. MVGALR. FLPWAPRGSKPIILEPRPFLSF RDFSEWSVHSGDSSPGLPEAANPLYWNLGPS. V	CCCCTCCACACACAGCTAGGAGCCCATGCCCGGCCTGATCTCAGCCCGAGGACAGCCCCTCCTTGAGGTCCTTCCT	PLHTARSPCPA, SQPEDSPSLRSFLPKPTWVPS PSTQLGAHARPDLSPRTAPP, GPSSPPGCPL SPPHS, EPMPGLISARGQPLLEVLPPQAHLGALF

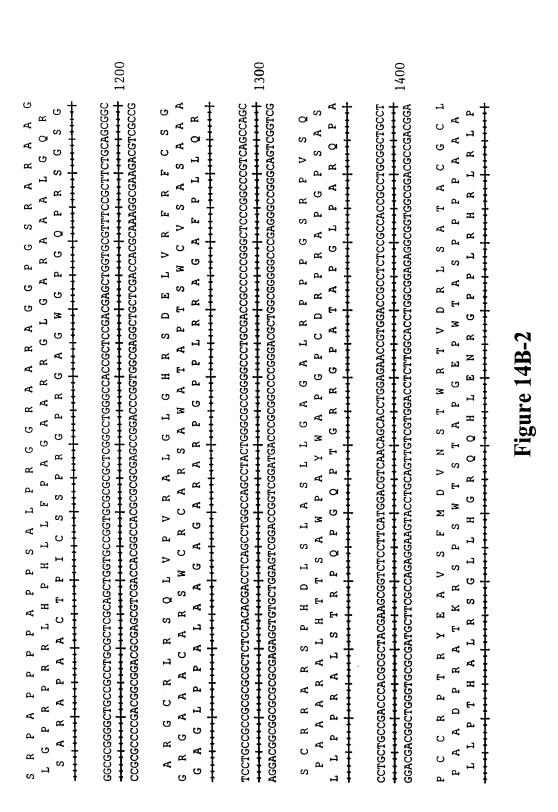
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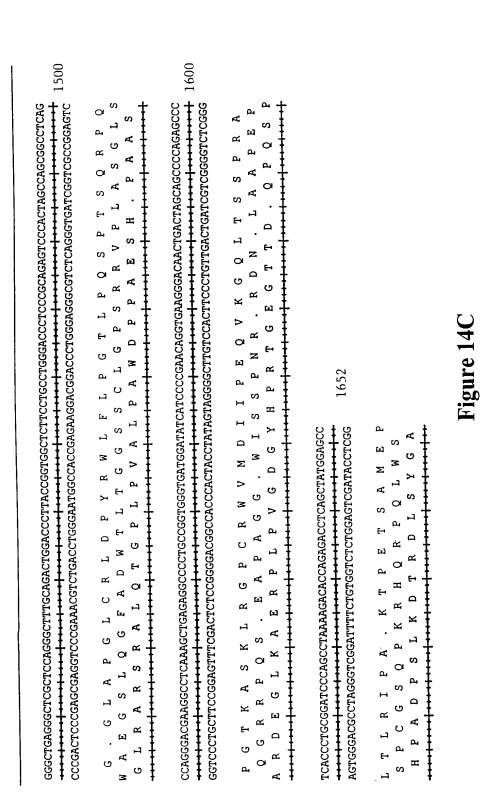
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38/40



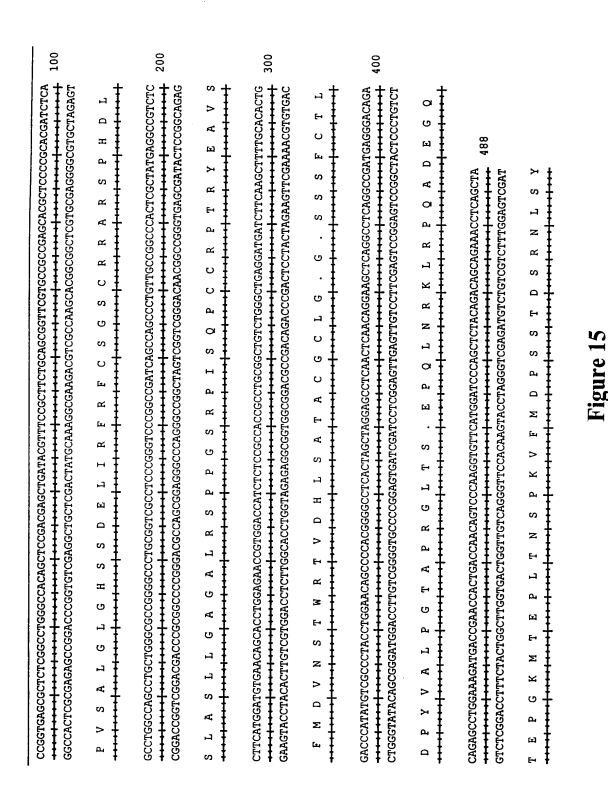
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#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/22604

A. CLASSIFICATION OF SUBJECT MATTER						
IPC(7) :Please See Extra Sheet. US CL :Please See Extra Sheet.						
According to International Patent Classification (IPC) or to both national classification and IPC						
B. FIELDS SEARCHED						
Minimum documentation searched (classification system follow	ved by classification symbols)					
U.S. : Please See Extra Sheet.						
Documentation searched other than minimum documentation to t	he extent that such documents are included in the fields searched					
Electronic data base consulted during the international search ( Please See Extra Sheet.	name of data base and, where practicable, search terms used)					
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category* Citation of document, with indication, where a	appropriate, of the relevant passages Relevant to claim N	о.				
A US 5,739,307 A (JOHNSON, JR (14/04/98), see entire document.	R. ET AL) 14 April 1998 1-38					
US 5,747,655 A (JOHNSON, JR. ET see entire document.	AL) 05 May 1998 (05/05/98), 1-38					
Further documents are listed in the continuation of Box	C. See patent family annex.	l				
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17 DECEMBER 1999	<b>10</b> FEB 2000					
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Washington, D.C. 20231	PREMA MERTZ					
Facsimile No. (703) 305-3230	Telephone No. (703) 308-0196 / /	- 1				

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/22604

A. CLASSIFICATION OF SUBJECT MATTER: IPC (7):

C07K 14/47, 14/475; C12N 5/10, 15/12, 15/16, 15/63, 15/64; A61K 38/16, 38/17, 38/18, 39/395, 48/00

A. CLASSIFICATION OF SUBJECT MATTER: US CL :

530/350,399; 536/23.1, 23.5, 23.51, 24.3, 24.31; 514/2, 8, 12, 44; 435/69.1, 69.4, 70.1, 71.1, 71.2, 471, 325, 375, 252.3, 254.11, 320.1, 6, 7.1, 7.2

#### **B. FIELDS SEARCHED**

Liminum documentation searched

Classification System: U.S.

530/350,399; 536/23.1, 23.5, 23.51, 24.3, 24.31; 514/2, 8, 17, 44, 42/69.1, 69.4, 70.1, 71.1, 71.2, 471, 325, 375, 252.3, 254.11, 320.1, 6, 7.1, 7.2

#### B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

WEST, CAS ONLINE, MEDLINE, EMBASE, BIOSIS

search terms: artemin, growth factor, polypeptide, polynucleotide, cDNA, GFR alpha, recombinant production, antibody, detection, assay, hybridization, treatment, method, therapy.